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Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)

McAleenan A, Kelly C, Spiga F, Kernohan A, Cheng HY, Dawson S, Schmidt L, Robinson T, Brandner S, Faulkner CL, Wragg C, Jefferies S, Howell A, Vale L, Higgins JPT, Kurian KM

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[Prognosis Review]

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide

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ABSTRACT

Background

Glioblastoma is an aggressive form of brain cancer. Approximately five in 100 people with glioblastoma survive for five years past diagnosis. Glioblastomas that have a particular modification to their DNA (called methylation) in a particular region (the O⁶-methylguanine–DNA methyltransferase (MGMT) promoter) respond better to treatment with chemotherapy using a drug called temozolomide.

Objectives

To determine which method for assessing MGMT methylation status best predicts overall survival in people diagnosed with glioblastoma who are treated with temozolomide.

Search methods

We searched MEDLINE, Embase, BIOSIS, Web of Science Conference Proceedings Citation Index to December 2018, and examined reference lists. For economic evaluation studies, we additionally searched NHS Economic Evaluation Database (EED) up to December 2014.

Selection criteria

Eligible studies were longitudinal (cohort) studies of adults with diagnosed glioblastoma treated with temozolomide with/without radiotherapy/surgery. Studies had to have related MGMT status in tumour tissue (assessed by one or more method) with overall survival and presented results as hazard ratios or with sufficient information (e.g. Kaplan-Meier curves) for us to estimate hazard ratios. We focused mainly on studies comparing two or more methods, and listed brief details of articles that examined a single method of measuring MGMT promoter methylation. We also sought economic evaluations conducted alongside trials, modelling studies and cost analysis.

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Data collection and analysis

Two review authors independently undertook all steps of the identification and data extraction process for multiple-method studies. We assessed risk of bias and applicability using our own modified and extended version of the QUality In Prognosis Studies (QUIPS) tool. We compared different techniques, exact promoter regions (5'-cytosine-phosphate-guanine-3' (CpG) sites) and thresholds for interpretation within studies by examining hazard ratios. We performed meta-analyses for comparisons of the three most commonly examined methods (immunohistochemistry (IHC), methylation-specific polymerase chain reaction (MSP) and pyrosequencing (PSQ)), with ratios of hazard ratios (RHR), using an imputed value of the correlation between results based on the same individuals.

Main results

We included 32 independent cohorts involving 3474 people that compared two or more methods. We found evidence that MSP (CpG sites 76 to 80 and 84 to 87) is more prognostic than IHC for MGMT protein at varying thresholds (RHR 1.31, 95% confidence interval (CI) 1.01 to 1.71). We also found evidence that PSQ is more prognostic than IHC for MGMT protein at various thresholds (RHR 1.36, 95% CI 1.01 to 1.84). The data suggest that PSQ (mainly at CpG sites 74 to 78, using various thresholds) is slightly more prognostic than MSP at sites 76 to 80 and 84 to 87 (RHR 1.14, 95% CI 0.87 to 1.48). Many variants of PSQ have been compared, although we did not see any strong and consistent messages from the results. Targeting multiple CpG sites is likely to be more prognostic than targeting just one. In addition, we identified and summarised 190 articles describing a single method for measuring MGMT promoter methylation status.

Authors' conclusions

PSQ and MSP appear more prognostic for overall survival than IHC. Strong evidence is not available to draw conclusions with confidence about the best CpG sites or thresholds for quantitative methods. MSP has been studied mainly for CpG sites 76 to 80 and 84 to 87 and PSQ at CpG sites ranging from 72 to 95. A threshold of 9% for CpG sites 74 to 78 performed better than higher thresholds of 28% or 29% in two of three good-quality studies making such comparisons.

PLAIN LANGUAGE SUMMARY

Which method of determining MGMT promoter methylation best predicts survival in people with glioblastoma treated with temozolomide?

What was the aim of this review?

Glioblastoma is a very aggressive type of brain cancer. People with glioblastoma are usually treated with surgical removal of the tumour followed by radiotherapy, chemotherapy, or both. The standard chemotherapy is a medicine called temozolomide. Some glioblastoma tumours have a particular modification in their DNA (which contains the genetic code of organisms), and knowing whether a person has this modification is useful to predict how long the person may live after their diagnosis with cancer and how they may respond to temozolomide. The modification is known as 'methylation of the MGMT promoter region' and it can also affect MGMT protein expression (the way MGMT is made and modified). There are many ways to work out whether a tumour has this modification. In this review, we attempted to work out which method is best.

What we found

We identified 32 studies comparing different ways to measure whether the MGMT promoter region is methylated. The main three methods were called 'methylation-specific polymerase chain reaction (PCR),' 'pyrosequencing' (both of which look directly at the MGMT promoter region) and 'immunohistochemistry' (which looks at MGMT protein expression). We found that methylation-specific PCR and pyrosequencing are better at predicting overall survival than immunohistochemistry. Methylation-specific PCR and pyrosequencing can be carried out by targeting different parts of the tumour DNA. Pyrosequencing can be performed using different cut-off thresholds to determine whether a tumour is methylated or unmethylated. We did not identify very clear signals in terms of the best parts of the DNA to target or which are the best cut-off thresholds.

How reliable are results of the studies in this review?

We rated our confidence in the evidence as 'moderate' for our conclusions about methylation-specific PCR, but as 'low' for pyrosequencing. Although there were many studies, they all looked at different variants of the methods, so it is difficult to work out exactly which variant is best.

What are the implications of this review?

Our review indicates both methylation-specific PCR and pyrosequencing provide better predictions of survival than immunohistochemistry. There is some evidence that pyrosequencing may be better than methylation-specific PCR at predicting overall survival, depending on the DNA targets and cut-off thresholds used. We documented the most frequent DNA targets used in methylation-specific PCR and pyrosequencing. We described cut-off thresholds used in pyrosequencing, although it is unclear which of these is best.



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SUMMARY OF FINDINGS

Summary of findings 1. Methods for measuring MGMT promoter methylation status

Methods for measuring MGMT promoter methylation status

Patient or population: people with glioblastoma undergoing treatment with temozolomide

Outcome being predicted: overall survival (time to death)

Technique/method	Ratio of hazard ra- tios (95% CI)	No of participants (studies)	Certainty of the evi- dence (GRADE)	Comments
MSP compared with IHC	1.31 (1.01 to 1.71)	913	$\oplus \oplus \oplus \odot$	_
		(7 studies)	Moderate ^a	
PSQ compared with IHC	1.36 (1.01 to 1.84)	871		_
		(5 studies)	Low ^b	
PSQ compared with MSP	1.14 (0.87 to 1.48)	1119	000	_
		(9 studies)	Low ^b	
Variants of PSQ	Not estimated	876	000	_
		(11 studies)	Very low ^c	
qMSP (against MSP or PSQ)	Not estimated	765	\$000	_
		(7 studies)	Very low ^c	
Bead array (against MSP or PSQ)	Not estimated	81	000	_
		(2 studies)	Very low ^d	
PCR-mRNA (against MSP or PSQ)	Not estimated	148	000	_
		(2 studies)	Very low ^e	
MS-MLPA (against MSP or PSQ)	Not estimated	48	000	_
		(1 study)	Very low ^f	
PCR-HRM (against MSP or PSQ)	Not estimated	309	000	_
		(3 studies)	Very low ^g	
Other techniques (against MSP	Not estimated	1209	000	_
or PSQ)		(7 studies) across various other techniques	Very low ^d	

Grades of evidence

High certainty: further research is very unlikely to change our confidence in the conclusion.

Moderate certainty: further research is likely to have an important impact on our confidence in the conclusion. **Low certainty:** further research is very likely to have an important impact on our confidence in the conclusion. **Very low certainty:** we are very uncertain about the conclusion.

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CpG: 5'-cytosine-phosphate-guanine-3'; **HRM:** high-resolution melting; **IHC:** immunohistochemistry; **MGMT:** O⁶-methylguanine–DNA methyltransferase; **MS-MLPA:** methylation-specific multiplex ligation-dependent probe amplification; **MSP:** methylation-specific polymerase chain reaction; **PCR:** polymerase chain reaction; **PCR:** polymerase chain reaction-messenger ribonucleic acid; **PSQ:** pyrosequencing; **qMSP:** quantitative methylation-specific polymerase chain reaction.

^aDowngraded one level for imprecision.

^bDowngraded two levels for imprecision and indirectness (due to variability in CpG sites and thresholds used for PSQ).

^cDowngraded three levels for serious risk of bias, imprecision, inconsistency and indirectness.

 $^{\rm d}\mbox{Downgraded}$ three levels for serious imprecision, inconsistency and indirectness.

^eDowngraded three levels for imprecision, inconsistency and indirectness.

^fDowngraded three levels for serious risk of bias, serious imprecision, inconsistency and indirectness.

gDowngraded three levels for risk of bias, serious imprecision, inconsistency and indirectness.



BACKGROUND

Description of the health condition and context

Glioblastoma is an aggressive form of brain cancer. Approximately five of every 100 people with glioblastoma survives for five years past diagnosis (Ostrom 2014). Glioblastomas that have a particular modification to their DNA (called methylation) in a particular region (the O⁶-methylguanine–DNA methyltransferase (MGMT) promoter) respond better to treatment with chemotherapy using a drug called temozolomide. Although we know that modification of this DNA region is important (Butler 2020), we do not know the best way to measure it. In this Cochrane Review, we aimed to assess which way of measuring methylation of the MGMT promoter best predicts survival for people with glioblastoma who are treated with temozolomide.

Gliomas are a group of brain tumours that share some features with glial cells, which are the cells that support and insulate neurons and are thought to originate from a population of stem or progenitor cells in the brain. The World Health Organization (WHO) divides gliomas into astrocytic, oligodendroglial and ependymal tumours, and other rarer subtypes depending on the type of glial cell the tumour shares features with (Louis 2016). Glioblastoma is the most malignant (aggressive) type of astrocytic tumour (Louis 2016), and the most common primary brain tumour among adults. Age-adjusted incidence of primary (isocitrate dehydrogenase (IDH)wild-type) glioblastoma (ICD-O-3 morphology codes 9440 to 9442, WHO grade IV) ranges from 0.59 to 3.69 per 100,000 people (Ostrom 2014). IDH-wild-type glioblastomas are more common in older people, peaking in 74 to 84-year olds (Ostrom 2014). These glioblastomas are associated with poor prognosis, with a five-year relative survival of approximately 5% (Ostrom 2014). The median overall survival is 9.9 months for people treated with surgery plus radiotherapy, and 15 months for people treated with surgery plus radiotherapy plus chemotherapy (Louis 2016). For people with secondary (IDH-mutant) glioblastomas, median overall survival is 24 months for people treated with surgery plus radiotherapy, and 31 months for people treated with surgery plus radiotherapy plus chemotherapy (Louis 2016).

Glioblastomas are commonly diagnosed by a neurosurgical multidisciplinary team following brain imaging with computerised tomography (CT) and magnetic resonance imaging (MRI). If appropriate, the person has a biopsy or resection (surgical removal) of the tumour to confirm the histopathological diagnosis. For newly diagnosed glioblastoma, the standard treatment is maximal surgical resection followed by radiotherapy with concomitant and adjuvant temozolomide (Stupp 2005). Temozolomide is an alkylating chemotherapeutic agent. It causes DNA damage, which inhibits DNA replication. However, not all people respond to temozolomide therapy to the same extent. There is evidence that people with newly diagnosed glioblastoma who start treatment with radiotherapy and temozolomide more than six weeks after neurosurgery have worse overall survival than people who start treatment within six weeks (Sun 2015).

In the UK, it is estimated that on average just over 20 years of life are lost per person with a brain tumour, the most of any form of cancer (Burnet 2005). Olesen 2012 estimated the total annual costs of brain tumours in Europe to be EUR 5.2 billion, based upon purchasing power parity rates for 2010.

Description of the prognostic factors

MGMT is a DNA repair enzyme in glioblastoma cells that can repair the damage caused by alkylating agents such as temozolomide. If the MGMT gene promoter is methylated, it is thought the glioblastoma cell is less able to repair this damage and is more likely to die, therefore making the tumour more sensitive to alkylating therapy (Brandner 2015). If the MGMT gene promoter in the glioblastoma cell is unmethylated, it is thought that the glioblastoma cell can repair the damage caused by temozolomide and, therefore, temozolomide is less effective. Consequently, epigenetic silencing of the MGMT gene by promoter methylation is associated with longer overall survival in people with glioblastoma receiving alkylating therapy in addition to radiotherapy (Alnahhas 2020; Esteller 2000; Hegi 2004; Hegi 2005). A key retrospective analysis of one randomised phase III trial found that treatment with temozolomide and radiotherapy conferred a significant survival benefit versus radiotherapy alone in people with MGMT promoter methylation (median survival: 21.7 months, 95% confidence interval (CI) 17.4 to 30.4 with temozolomide plus radiotherapy versus 15.3 months, 95% CI 13.0 to 20.9 with radiotherapy alone; P = 0.007), whereas there was a smaller difference in survival in people with unmethylated MGMT (median survival: 12.7 months, 95% CI 11.6 to 14.4 with temozolomide plus radiotherapy versus 11.8 months, 95% CI 9.7 to 14.1 with radiotherapy alone) (Hegi 2005).

There is clear evidence that MGMT promoter methylation status testing is important in older people. When older people with glioblastomas with an unmethylated MGMT promoter were treated with single-agent temozolomide chemotherapy, they had worse outcomes than those treated with radiotherapy (Malmström 2012; Wick 2012). Professional bodies, such as the European Association for Neuro-Oncology (EANO), recommend evaluation of MGMT promoter methylation status in older people (Weller 2017a). The National Institute for Health and Care Excellence (NICE) recommends that all high-grade gliomas are tested for MGMT promoter methylation to inform prognosis and guide treatment (NICE 2018). Most non-elderly (aged under 65 years) people are treated with temozolomide chemotherapy irrespective of MGMT promoter status, possibly due to the lack of alternative treatments (Hegi 2015). Despite this, MGMT promoter status is still a useful prognostic marker which may impact clinical management. It can also inform recruitment into clinical trials for novel therapies.

There are many ways of assessing methylation status. These include:

- methylation-specific polymerase chain reaction (MSP);
- quantitative (real-time) methylation-specific polymerase chain reaction (qMSP), including MethyLight;
- methylation-specific sequencing, including pyrosequencing (PSQ);
- bead array;
- methylation-specific multiplex ligation-dependent probe amplification (MS-MLPA);
- polymerase chain reaction with high-resolution melting (PCR-HRM);
- co-amplification at lower denaturation temperature (COLD)-PCR; and
- digestion-based assays.

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We describe these techniques briefly in Table 1. In addition, protein expression or enzymatic activity may be used as a proxy for methylation status. Internationally accepted consensus about the most appropriate diagnostic method for MGMT promoter status is lacking (Brandner 2015). MSP was used to assess MGMT promoter status in the landmark study by Hegi 2005. In practice, the choice of technique used to assess MGMT promoter status may depend on the amount and quality of the DNA sample(s) (e.g. formalinfixed paraffin-embedded (FFPE) versus frozen tissue-derived DNA), the robustness and simplicity of the method, the availability of equipment and reagents, cost and experience. In the most recent UK National Quality Assessment (UK NEQAS) External Quality Assessment report, 10 of 18 UK laboratories used PSQ, five used MSP, two used HRM and one used MS-MLPA.

Additionally, there is within-technique heterogeneity arising from differences in the regions of the MGMT promoter tested to determine MGMT methylation status. The prognostic impact of these differences is not well understood. Similarly, there is variation in the cut-offs used for categorising methylation status for techniques that quantify the amount of methylation present. The manufacturers do not recommend specific thresholds and there is no consensus on what are the most appropriate, with individual laboratories left to determine their own thresholds, for example by running tests on healthy control samples, or examining survival of people with glioblastoma.

The result of each method for measuring MGMT status can be considered a separate prognostic factor for predicting overall survival in people with glioblastoma treated with temozolomide.

Health outcomes

The health outcome of interest for this review was overall survival. We did not limit the period of follow-up. Glioblastomas are associated with poor prognosis, so we anticipated that most studies would assess overall survival within five years of diagnosis.

Why it is important to do this review

It is important to reach a consensus regarding which is the best method for assessing MGMT methylation status based on the prognostic value of each method in predicting overall survival in people with glioblastoma treated with temozolomide, so that people living with glioblastoma can be confident that they are having the appropriate molecular analysis performed. The regions of the promoter that need to be analysed and the most relevant cut-offs for quantitative tests need to be established. Systematic reviews and meta-analyses have determined the prognostic value of MGMT promoter status assessed by a specific technique, for example by PSQ (Zhao 2016), MSP (Zhang 2013), or qMSP (Hegi 2019). However, we are aware of no systematic review that has determined which method is best correlated with prognosis. One narrative overview addressed the question, but provided no quantitative synthesis of the results (Dullea 2016).

In this Cochrane Review, we seek to determine which technique, assessing which regions and (if relevant) which cut-off is best associated with overall survival in people with glioblastoma treated with temozolomide. We consider each MGMT test as a separate prognostic factor. We extract or calculate (where possible) hazard ratios (HRs) for those who tested positive compared with those who tested negative. A test that is not better than flipping a coin is expected to have an HR of one. The better the test's ability to discriminate between people with a good overall survival versus people with poor overall survival, the further the HR value will be from one.

The review aims to answer part of the question "Do molecular subtyping techniques improve treatment selection, prediction and prognostication in people with brain and spinal cord tumours," one of the top 10 topics identified by the James Lind Alliance Neuro-Oncology Priority Setting Partnership (JLA PSP 2018), by addressing the predictive ability of one specific molecular modification (MGMT methylation status) in people with glioblastoma. The James Lind Alliance is an organisation that brings people, carers and clinicians together to set research priorities. The National Cancer Research Institute Brain Tumour Clinical Studies Group has also identified this as an area for future research.

It is also important to consider the cost effectiveness of alternative methods of assessing MGMT promoter methylation status. Each method of assessment will incur costs, such as laboratory costs, clinic costs and subsequent treatment costs. The benefits of targeting treatment may include greater survival and less exposure to potentially toxic treatments, as well as potential cost-savings from the avoidance of waste from the use of ineffective drugs. This review aims to consider the costs alongside the consequences of the prognostic tests to understand the value that they provide to the healthcare system.

OBJECTIVES

Primary objective

To determine which method for assessing MGMT methylation status best predicts overall survival in people diagnosed with glioblastoma who are treated with temozolomide. We consider each MGMT method as a separate prognostic factor.

See Table 2 for the review question in population, index prognostic factor, comparator prognostic factor(s), outcome, timing and setting (PICOTS) format.

Secondary objective

We undertake an integrated economic review to identify economic evaluations in relation to the different methods of assessing MGMT methylation status effect on overall survival, and undertake a simple economic analysis exploring the cost-effectiveness of alternative approaches to assessing MGMT methylation status.

Investigation of sources of heterogeneity

We examine for each technique whether any of the following features was best associated with overall survival.

- Promoter region/CpGs analysed (or the antibody used in the case of IHC).
- Cut-off used (where relevant).
- Type of tumour sample (FFPE or frozen).

We planned to investigate the effect of population characteristics including the following if sufficient data allowed us to do this.

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• Age.

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- Extent of tumour resection.
- Karnofsky performance status.
- IDH status.
- Recurrent tumour versus first diagnosis.

We are assuming constant HRs. To confirm the validity of this assumption, we hoped to investigate length of follow-up as a source of heterogeneity, again if sufficient replications of the same methods had been available.

METHODS

Criteria for considering studies in this review

Types of studies

We included longitudinal studies of adults with diagnosed glioblastoma treated with temozolomide with/without radiotherapy/surgery that had related MGMT status in tumour tissue (assessed by one or more method) with overall survival. This included the temozolomide-treated arms of randomised controlled trials (RCT). We also sought nested case-control studies. To be included, studies must have determined MGMT status from samples taken before the initiation of treatment. Studies could have had any length of follow-up. We excluded cohort studies performed exclusively in people who had survived a particular amount of time, or case reports.

Studies were only eligible if they reported HRs, or if we could calculate HRs from the data reported.

Types of studies for the economic component

We sought economic evaluations conducted alongside trials, modelling studies and cost analyses to inform the identification of cost-effectiveness outcomes.

Targeted participants

Eligible studies were of adults with diagnosed glioblastoma treated with temozolomide with or without radiotherapy/surgery. If studies included people with other forms of glioma (and we could not extract results for the population with glioblastoma), we included these if other forms of glioma made up less than 10% of the population. We included studies of participants with either first diagnosis or recurrent glioblastoma. Participants in eligible studies could receive concomitant and adjuvant therapies in addition to temozolomide (e.g. surgery or radiotherapy, or both, or additional chemotherapeutics). If not all participants received temozolomide (e.g. in the context of an RCT), we included data on people who did receive temozolomide if these were available. We excluded studies performed exclusively in children (under 18 years of age).

Types of prognostic factors

Eligible studies had to assess MGMT promoter methylation status in tumour tissue by at least one method. We treated each method as a separate prognostic factor. Eligible techniques included, but were not restricted to, MSP; quantitative MSP (real-time PCR or MethyLight methylation-specific quantitative PCR); methylationspecific sequencing, including PSQ; bead array; MS-MLPA; PCR-HRM; COLD-PCR and digestion-based assays. We also included testing strategies that considered MGMT expression (e.g. IHC for protein expression, or tests measuring messenger ribonucleic acid (mRNA) levels) or MGMT enzymatic activity. Eligible techniques had to be molecular techniques and performed directly on tumour tissue. We excluded studies that assessed MGMT promoter methylation status from blood samples because insufficient quantities of brain tumour DNA cross the blood-brain barrier for testing to be appropriate. In addition, we excluded studies that inferred MGMT methylation status due to macroscopic morphological changes that can be detected by, for example, imaging (i.e. MRI, CT, positron emission tomography (PET)).

We excluded studies that did not report the method of determining MGMT promoter methylation status, as this information is essential for this review.

Types of outcome to be predicted

• Overall survival.

Outcomes of the economic component

- Resources use, costs, cost effectiveness and cost-utility of different methods of assessing MGMT promoter methylation status based on full economic review.
- Relative efficiency of each method of testing for MGMT promoter methylation status based on a decision model using the outcomes from the review of effectiveness and from the full integrated economic review.

Search methods for identification of studies

Electronic searches

We searched the following databases in December 2018 (Appendix 1):

- Ovid MEDLINE (1946 to 4 December 2018);
- PubMed NOT MEDLINE (4 December 2018);
- Ovid Embase (1980 to 2018, week 49);
- BIOSIS (1969 to 3 December 2018) and
- Web of Science Conference Proceedings Citation Index (CPCI-S) (1900 to 3 December 2018).

We applied no restrictions on language or date of publication to the searches.

Searching other resources

The Society of Neuro-Oncology (SNO), and its partner associations the EANO and the Japan Society of Neuro-Oncology hold meetings where relevant research may be presented. We searched for abstracts from these meetings and other relevant conferences via the Web of Science Conference Proceedings Citation Index (CPCI-S) (from 1990 to 3 December 2018), as listed above. We translated the BIOSIS search for CPCI-S, since both databases are hosted on Web of Science.

Additional searches for the economic component

We searched the NHS Economic Evaluation Database (EED), with combinations of relevant keywords from the search strategy, up to the end of December 2014, when the last records were added to that database. The NHS EED was based on a comprehensive search of bibliographic databases including MEDLINE and Embase.

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Data collection

We used EPPI-Reviewer 4 and EPPI-Reviewer Web for the screening and selection of studies, and for select data extraction tasks (EPPI-Reviewer). Further data extraction was undertaken into a Microsoft Excel spreadsheet.

Selection of studies

Two review authors (of AM, KMK and AH) independently screened titles and abstracts of all identified search results. We retrieved the full text of any articles that either review author deemed relevant, or whose relevance could not be determined from the abstract. Two review authors (of AM, CK, FS, KMK, AH, SB and CLF) independently assessed the full-text articles for eligibility. We resolved any disagreements about eligibility as multiple-method studies by consensus, or by consulting a third review author where necessary. One review author made decisions about studies considered eligible as single-method studies. We constructed a PRISMA flow diagram to depict the flow of information through the different phases of the review.

Two review authors (AK and TR) screened studies retrieved for full-text screening for potentially relevant economic studies.

Data extraction and management

As planned in the protocol (McAleenan 2019), we performed full data extraction, risk of bias assessment and synthesis on studies that evaluated MGMT promoter methylation status of the same people using two or more methods (i.e. multiple techniques, CpG sites, cut-offs or types of tissue sample) so that these tests could be compared on the same samples of people. We performed limited data extraction on studies that evaluated MGMT promoter methylation status using a single method.

Two review authors (of AM, CK, FS, LS, HC and JPTH) independently performed data extraction on each article describing two or more methods for MGMT promoter methylation status, using forms piloted on several articles. We resolved any disagreements by consensus and consulted a third review author where necessary. We extracted data on the following items relevant to prognostic factor studies, derived from the CHecklist for critical Appraisal and data extraction for systematic Reviews of prediction Modelling Studies (CHARMS) (Moons 2014). We did not contact primary investigators for information that was not available in the reports.

Study characteristics

- Author.
- Year.
- Country and setting.
- Length of follow-up.
- Study dates.
- Study design.

Population characteristics

- Number of participants.
- Population source and setting.
- Timing of MGMT promoter methylation assessment.
- Inclusion/exclusion criteria.
- Tumour type.

- Age.
- Gender.
- Karnofsky performance status.
- Extent of resection.
- Treatment regimen.
- Length of time between neurosurgery and start of treatment.
- IDH mutation status.
- First diagnosis or recurrent disease.
- Deaths during follow-up.
- Prevalence of MGMT promoter methylation (by each technique).

Method(s) of MGMT promoter methylation assessment

- Technique.
- Tumour sample type (i.e. FFPE or frozen tissue).
- Region/CpGs analysed (for PCR-based tests); antibody used (for immunohistochemistry).
- Cut-off/threshold used to determine MGMT promoter methylation status (where relevant).
- Method of determining threshold and whether it was prespecified.

Outcome assessment

• Time point from which overall survival was measured.

Missing data

• Number of participants with any missing data.

Association between MGMT methylation status and overall survival

- Data sufficient to determine computer HRs and their Cls.
- Adjusted HRs and their CIs (where reported), and factors for which the result was adjusted.

To obtain HRs, we followed strategies described by Tierney 2007 and Parmar 1998. We primarily sought unadjusted HRs, and used these if they were presented directly. We computed standard errors of log HRs from CIs or exact P values, assuming these were based on Wald tests. Where only a P value threshold was stated, we set the P value to be equal to this; this only occurred in cases where small thresholds had been used (P < 0.001, or < 0.000001). When we could not obtain unadjusted HRs directly using these approaches, we obtained HRs using (in order of preference): 1. individual participant data (IPD) from publications; 2. reported adjusted HRs or 3. published Kaplan-Meier curves. From Kaplan-Meier curves, we reconstructed approximate IPD following Guyot 2012. We derived plot co-ordinates from the published curves using Engauge Digitizer 12.1 as input into Guyot's algorithm (Engauge Digitizer). Where possible, we followed Guyot and colleagues' suggestion of including information from risk tables and total numbers of events. Depending on the information provided in study reports, we reconstructed IPD using the best information (i.e. in preferential order 'all information,' 'no numbers at risk' then 'no total events' as referred to by Guyot 2012). However, for most study reports there was insufficient information, in which case we followed Guyot and colleagues' 'neither' case. We reconstructed the IPD using the R script from the supplement of Guyot 2012. These analyses were conducted using R (version 4.0.3) in RStudio (version

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1.2.5042). Reconstructed data from these plots are available at the data.bris repository (data.bris.ac.uk/data/).

We analysed the IPD or reconstructed IPD (from Kaplan-Meier curves) to estimate HRs using Cox proportional hazards regression, using the *stcox* command in Stata. Some study reports categorised participants by the extent of methylation, e.g. "unmethylated (0 to 9%)," "weakly methylated (10% to 29%)," and "methylated (30% or greater)." Where survival data for these groups were presented in Kaplan-Meier curves, we combined the individuals across categories to dichotomise the data at each cut-off. To illustrate, for the categories in the above example, we regrouped the data to estimate the HR for the comparison "unmethylated" and "weakly methylated" combined versus "methylated" (cut-off at 29%), and the HR for the comparison "unmethylated" versus "weakly methylated" and "methylated (cut-off 9%). These analyses, including plotting of reconstructed Kaplan-Meier curves, were performed using Stata (version 16).

For studies that evaluated MGMT promoter status using a single method, we extracted details on author, year, country, length of follow-up, number of participants, tumour type, IDH mutation status and technique used for MGMT promoter methylation assessment.

Economic studies

In addition to the data extracted from clinical studies, we planned to extract relevant data from economic evaluations (had any been identified). We aimed to collect the following data from the economic evaluation studies.

- Type of evaluations.
- Sources of effectiveness data.
- Cost data.
- Sources of cost data.
- Sources of outcome valuations.
- Analytical approach.

Assessment of risk of bias in included studies

We assessed risk of bias in studies that evaluated MGMT promoter methylation status of the same people using at least two methods.

The QUality In Prognosis Studies (QUIPS) tool is designed to assess risk of bias in prognostic factor studies (Hayden 2013). It assesses bias across six domains: study participation, study attrition, prognostic factor measurement, outcome measurement, study confounding, and statistical analysis and reporting. We assessed risk of bias across QUIPS domains, although we added a domain on subsequent treatment. We renamed the study confounding domain to 'adjustment for other potential prognostic factors'; and we limited the domain about statistical analysis and reporting to selective reporting alone because we sought only results of a standard proportional hazards regression analysis. We replaced the prompting items and considerations, which mainly assessed reporting, with signalling questions to help us reach domain-level judgements. The domain modifications and signalling questions were informed by the CHARMS checklist (Moons 2014), a framework for assessing internal validity of articles dealing with prognosis described in Altman 2001, as well as ROBINS-I (risk of bias in non-randomised studies of interventions) (Sterne 2016) and QUADAS-2 (Whiting 2011). In addition, for each domain apart from study attrition and selective reporting, we added questions assessing the applicability of the study as in QUADAS-2 (Whiting 2011) and PROBAST (Wolff 2019). We assessed risk of bias in the first three domains (participant selection, subsequent treatment and outcome measurement) at the study level, and the other four domains (prognostic factor measurement, study attrition, adjustment for other potential prognostic factors and selective reporting) for each result within each study. We judged risk of bias and concerns regarding applicability as high, low or unclear. The tool is detailed in Appendix 2. Two review authors (of AM, CK, FS, LS, HC and JPTH) independently performed assessments using a form that had been piloted on several articles. These review authors sought to reach a consensus judgement and resolved any remaining disagreements by consulting a third review author. We did not contact primary investigators for information that was not available in the reports.

Assessment of risk of bias in studies included in the economic component

We planned to perform assessment of the quality of the economic evaluations captured in this review in two stages. The first stage was to assess the risk of bias in the clinical studies informing the evaluation. If the economic evaluation was carried out alongside a single study, then we planned to use our bespoke tool described in Appendix 2. Should any economic evaluations based on models have been identified, we planned to assess any summary effect sizes from systematic reviews used as data inputs in these modelbased economic evaluations using the ROBIS tool (Whiting 2016). The second stage for assessing any identified economic evaluations was to assess the overall methodological quality of the economic component of the evaluation. Based on the methods from the Cochrane Handbook chapter on economic methods (Aluko 2020), we planned to assess evaluations carried out alongside single empirical studies using the CHEERS checklist (Husereau 2013). In addition, we planned to assess any model-based economic evaluations using the NICE methodology checklist (NICE 2012).

Assessment of reporting bias

For each meta-analysis that contained 10 or more studies, we planned to examine the symmetry of funnel plots and test for asymmetry using Debray's funnel inverse variance test based on HRs (Debray 2018). Asymmetry may be an indicator of publication bias.

Data synthesis

Data synthesis and meta-analysis approaches

To assess the relative prognostic ability of the different methods we focused on data from direct, within-study comparisons, where the MGMT promoter methylation status of the same series of people was evaluated in multiple ways and the results correlated with overall survival. We undertook full data extraction, risk of bias assessment and synthesis on studies only for this subset of studies.

We harmonised the direction of the HRs from each study so that each represented hazard rate among people with an unmethylated MGMT promoter divided by the hazard rate among people with a methylated MGMT promoter. This means that a value greater than one indicates favourable outcomes in people with a methylated MGMT promoter. The greater the HR, the better the method was at predicting time to death. In the main analyses focusing on

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unadjusted HRs, we substituted an adjusted HR if an unadjusted HR was not available. We present each statistical result with a 95% CI.

Where at least five studies had compared the same pair of methods, we compared the HRs within studies to produce a ratio of hazard ratios (RHR). A complication here was how to account for the correlation between the log HRs to reflect that the different methods were applied to the same people. We computed the correlation between the original test results from studies for which we could extract IPD from the publications. We assumed this correlation would carry approximately through to the HRs comparing the two methods. We then computed log HRs and their variances (the latter as var(logHR1) + var(logHR2) - 2Cov(logHR1, logHR2), with covariances computed from the imputed correlation coefficient). We performed standard random-effects meta-analyses (with DerSimonian-Laird estimator of between-study variance) to estimate an overall RHR. We performed a sensitivity analysis using higher and lower values for the correlation coefficient. In these analyses, we quantified heterogeneity across results of the studies using an estimate of the between-study variance in logRHRs and portrayed these using prediction intervals. We also reported between-study variance (Tau²). In addition, we describe the extent of inconsistency in the findings using the I^2 statistic, which describes the percentage of variation across studies that is due to heterogeneity rather than chance (Higgins 2002).

The prognostic value of each test may be dependent on other prognostic factors of overall survival, and these may have been adjusted for. In addition to analyses of unadjusted HRs, we planned to extract and meta-analyse adjusted results, to confirm that the tests have added prognostic value in addition to (easier to measure) prognostic factors such as age, gender, disease stage at diagnosis and comorbidity. We present HRs adjusting for age and extent of resection, which were the most common factors adjusted for.

We also expected to identify studies that had evaluated MGMT promoter using only one method. We present only brief details of these studies. At a later date we may investigate these studies further to supplement inferences from the comparative studies. Specifically, there may be a possibility of comparing methods indirectly across studies. Such unadjusted indirect comparisons rely on the assumption that the studies assessing each test for MGMT promoter methylation are similar for all important characteristics (i.e. that they were conducted on similar populations that had been given similar treatments (or that these factors were adjusted for) and that the risk of bias was similar). This is a very strong assumption, and not one we were willing to make in this review.

We planned to present the results from the full economic review as a narrative analysis, describing the results of the economic evaluations identified by the search. In addition to the narrative summary of the economic evaluations, we planned to use both the clinical and economic outcomes to inform a decision model to estimate the cost effectiveness of assessing MGMT status in the management of glioma.

Subgroup analysis and investigations of heterogeneity

We aimed to investigate potential sources of heterogeneity in the results for each method using subgroup analyses or metaregression, depending on the number of studies identified and the nature of the source of heterogeneity. We examined, for each technique, whether any of the following features was best associated with overall survival.

- The promoter region/CpGs analysed (or the antibody used in the case of immunohistochemistry).
- The cut-off used (where relevant).
- The type of tumour sample (FFPE or frozen).

We also planned to investigate the effect of population characteristics including:

- age;
- extent of tumour resection;
- Karnofsky performance status;
- IDH status;
- recurrent versus first diagnosis.

We assumed constant HRs. To test the validity of this assumption, we planned to investigate length of follow-up as a source of heterogeneity, and if studies had started follow-up for overall survival from different time points, we aimed to investigate this as a source of heterogeneity.

Sensitivity analyses

We planned sensitivity analyses restricting the analysis to studies at low or unclear risk of bias. We also performed sensitivity analyses imputing different correlation coefficients between logHRs within studies, as described in the section on 'Data synthesis and metaanalysis approaches.'

Decision model

We aimed to create an economic model using outcomes from both the clinical and economic evidence we identified. The aim was to use the extracted data to populate a decision analytic model, to assess the cost-effectiveness of different methods of testing for MGMT promoter methylation status in people with glioblastoma. The effect of the different methods of assessing MGMT promoter methylation status (including not assessing for promoter methylation status at all) was to be compared in terms of probability of effectiveness and overall survival. The model was to be conducted from the UK National Health Service perspective for a target population aged 65 years or over. The time horizon of the model in terms of costs considered would have been six weeks (i.e. until the start of temozolomide treatment). Key uncertainties were to be explored using sensitivity analysis. However, due to the paucity of evidence with which to parameterise a cost-effectiveness decision model, particularly in reliable estimates of costs, this was not possible.

As an alternative, we considered cost comparison ratios (CCRs) of the three main techniques (PSQ, MSP and IHC). The principle underpinning a CCR comes from the conditions required for an efficient allocation of resources. Economic theory determines that when resources are efficiently allocated, the ratio of marginal costs (MC) to marginal benefits (MB) for all treatments 'a' to 'n' must be equal (i.e. $MC_a/MB_a = MC_b/MB_b = MC_c/MB_c = MC_n/MB_n$). Rearranging this equation and simplifying shows that when allocation of resources is efficient, the ratio of marginal costs is equal to the ratio of marginal benefits of all care, such that MC_a/MC_b

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= MB_a/MB_b . To inform these analyses, we used costs for performing these tests in the UK.

Summary of findings

We present the prognostic value of each method on overall survival in a 'Summary of findings' table. We assessed confidence in each result using the GRADE approach (Guyatt 2008). Guidance on the use of GRADE for prognostic factor studies has not yet been published, although adaptations have been proposed (Huguet 2013). We rated the overall strength of evidence as 'high,' 'moderate,' 'low' or 'very low.' We considered risk of bias, indirectness, inconsistency, imprecision and publication bias, which may lead to downgrading of the strength of the evidence; and size of effect, which may lead to upgrading of the strength of the evidence (see Appendix 3).

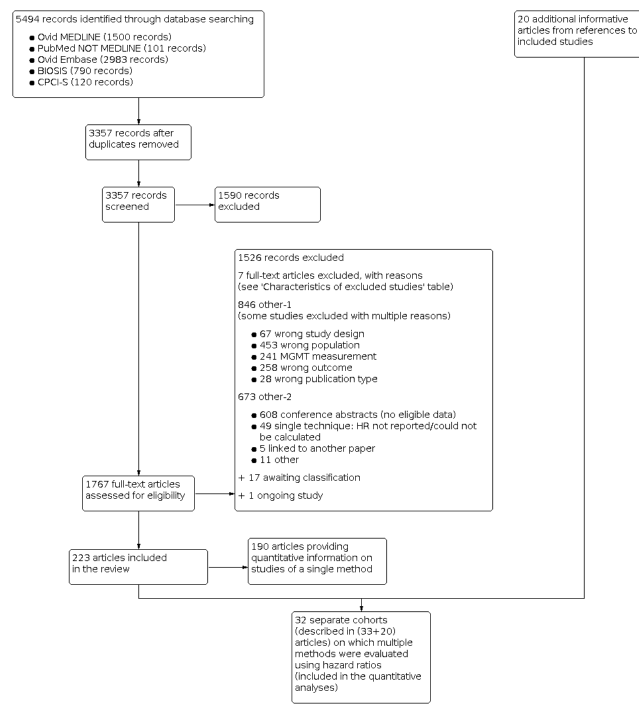
RESULTS

Results of the search

The search identified 5494 records and we included 223 of these in the review (see Figure 1).



Figure 1. Study flow diagram. N, number of records.



We included 32 distinct cohorts of people (referred to hereafter as studies) in our main analysis of studies comparing two or more methods, including studies comparing different variants of the same technique. These were reported in 33 articles. We drew on an additional 20 publications that had been cited in these 33 articles; some cohorts were reported in multiple publications, and some publications reported on two studies. In addition, we included 190 articles describing single-method studies. We summarise these in less detail than the main (multiple-method) studies, as planned in the protocol (McAleenan 2019).

None of the papers examined in full text met the criteria for inclusion into the review of economic evidence. The search of the NHS EED database also identified no relevant economic evaluations. Thus, we identified no economic evaluations assessing the cost-effectiveness of tests for MGMT promoter methylation status people with glioblastoma treated with temozolomide. This was true for evaluations based on a single study or using decision modelling to synthesise data from multiple studies. Thus, there is a lack of evidence for determining the most efficient strategies for assessing tests for MGMT promoter methylation status of people with glioblastoma treated with temozolomide.

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Characteristics of the included studies

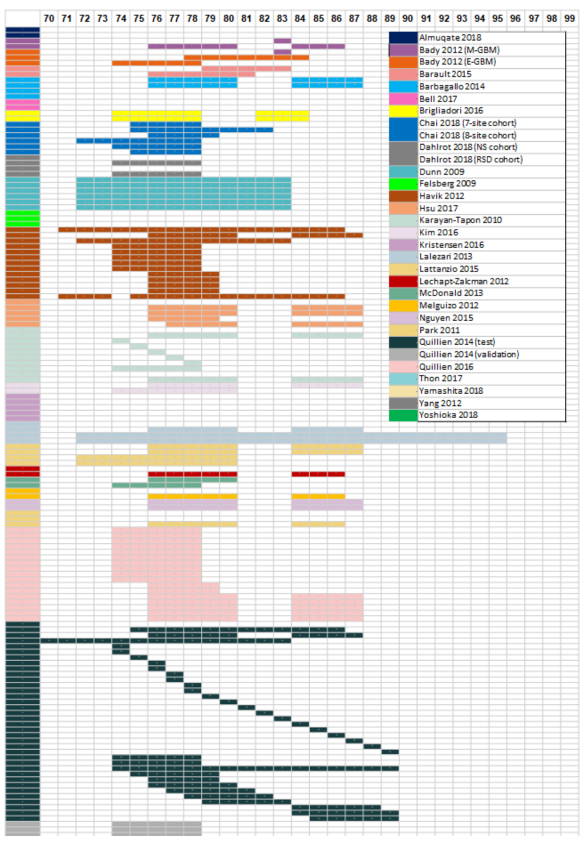
Details of the 32 studies, including 3474 participants, are presented in the Characteristics of included studies table. In descending order, the techniques investigated in the most studies were PSQ in 20 studies; MSP in 17 studies, IHC in nine studies, quantitative MSP (qMSP) in eight studies (including semiquantitative polymerase chain reaction (SQ-PCR), fluorescent semi-quantitative methylation-specific polymerase chain reaction (FSQ-MS-PCR) and MethyLight-MSP), PCR-HRM in three studies (Havik 2012; Quillien 2014 (test); Yamashita 2018); bead array in two studies (Bady 2012 (M-GBM); Bady 2012 (E-GBM)); polymerase chain reaction-targeting messenger ribonucleic acid (PCR-mRNA) in two studies (Felsberg 2009; Karayan-Tapon 2010); and MS-MLPA in one study (Park 2011). Other techniques studied were methylationspecific restriction enzyme quantitative polymerase chain reaction (MS-RE-qPCR; Almuqate 2018), methyl-beaming (Barault 2015), quantitative fluorescence immunohistochemistry (QF-IHC AQUA; Bell 2017), double immunofluorescence (DIF; Dahlrot 2018 (NS cohort); Dahlrot 2018 (RSD cohort)) qMSP combined with PSQ (qMSP-PSQ; Kristensen 2016), and sequencing (Thon 2017). The largest study compared qMSP with QF-IHC AQUA in 452 tumour samples (Bell 2017). The second largest compared MSP, PSQ and IHC in 418 samples (Lalezari 2013), and the third largest compared PSQ against DIF in 234 samples (Dahlrot 2018 (RSD cohort)). All other studies included fewer than 160 samples, with the smallest including 18.

All studies had a standard cohort design (with one being embedded in a randomised trial; Bell 2017). Nineteen studies were undertaken in Europe, two in North America, eight in East Asia, one in Australia and two across multiple countries. Mean ages ranged from 44 to 64 years. All studies had more men than women (overall, 60% were men where reported). Most studies were exclusively in people with glioblastoma at first diagnosis (where reported). In most studies, the majority of participants had undergone total resection. In 10 studies, it was explicitly stated that treatment followed the Stupp protocol (Stupp 2005), and in most of the others it was clear that temozolomide and radiotherapy were provided in a way that appeared consistent with the Stupp protocol.

We illustrate the comparisons made in the different studies in Table 3 and Appendix 4. Details of the specific methods implemented are provided in Appendix 5. We illustrate the CpG sites targeted in Figure 2.

Figure 2. Graphical illustration of the CpG sites examined. The top row indicates the number of the CpG site. Each row is colour-coded, corresponding to the enclosed legend indicating the study ID. Rows with blank cells (i.e. no colour-coded CPG sites) indicate that a method was not PCR-based test or that CpG information is not available. For

studies using PCR primers as described by Esteller 1999, CpG sites location is based on Bienkowski 2015. CpG: 5'cytosine-phosphate-guanine-3'; GBM: glioblastoma; NS: Nordic Study; RSD: Region of Southern Denmark.



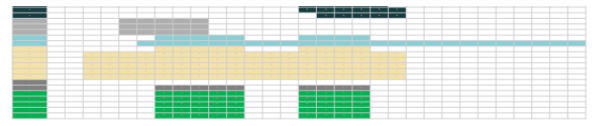
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Figure 2. (Continued)



Risk of bias assessment of included studies

We present results of the risk of bias assessment for the three domains that apply to the whole studies in Figure 3. All studies were assessed at low or unclear risk of bias for participant selection (domain D1). All studies except one were assessed to be at low risk of bias arising from variation in subsequent treatment after collection of the tumour sample (D2). All studies were assessed to be at low risk of bias in measurement of the outcome (all-cause mortality; D3). We present result-level risk of bias judgements in forest plots of these results in subsequent sections. We were mostly

free of concerns about risk of bias in the domains for study attrition (D5), problems with other prognostic factors adjusted for (D6) and selective reporting (D7). One PSQ result from Lalezari 2013 was deemed to be at high risk of bias due to attrition because more than 25% of the sample was missing from the analysis. Results for PCR-HRM from Yamashita 2018 were considered to be at high risk of bias from selective reporting because two other primer sets were used; one was discarded because of its inferior predictive performance. Full details of the risk of bias assessments and their justifications are available in Appendix 6.

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Figure 3. Study-level risk of bias assessments for studies comparing two or more methods. D1 = participant selection; D2 = subsequent treatment; D3 = outcome measurement. Green (+) = low risk of bias; yellow (-) = unclear risk of bias. GBM: glioblastoma; NS: Nordic Study; RSD: Region of Southern Denmark.

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Bady 2012 (E-GBM) • • • Bady 2012 (M-GBM) - • • Barault 2015 • • • Barbagallo 2014 • • • Barbagallo 2014 • • • Bell 2017 • • • Bell 2017 • • • Brigliadori 2016 - • • Chai 2018 (7-site cohort) - • • Chai 2018 (8-site cohort) - • • Dahlrot 2018 (RSD cohort) • • • Dunn 2009 • • • • Havik 2012 • • • • Kirayan-Tapon 2010 • • • • Kistensen 2016 • •			D1	D2	D3
Bady 2012 (M-GBM) - • • Barault 2015 • • • • Barbagallo 2014 • • • • Barbagallo 2014 • • • • Bell 2017 • • • • Bell 2017 • • • • Barbagallo 2016 - • • • Chai 2018 (7-site cohort) - • • • Chai 2018 (8-site cohort) - • • • • Dahlrot 2018 (RSD cohort) • • • • • • Dunn 2009 • • • • • • • Havik 2012 • • • • • • • Havik 2017 - • • • • • • Karayan-Tapon 2010 • • • • • • <td< td=""><td></td><td>Almuqate 2018</td><td>-</td><td>•</td><td>•</td></td<>		Almuqate 2018	-	•	•
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Our assessment of the applicability of the studies to the typical clinical context found almost all the studies to be generally applicable. We had no concerns for any studies regarding applicability issues in domains of subsequent treatment (D2) or prognostic factor measurement (D4). All studies except one were free of applicability concerns in the domains of outcome measurement (D3) and other prognostic factors adjusted for (D6); in these exceptional studies we were unclear rather than concerned. In the final domain of assessment for applicability (participant selection), we had high concerns for just one study, which included only people who were treated with open resection and who received at least two cycles of chemotherapy, thus excluding people of poorer clinical condition and those who did not survive to complete two cycles of chemotherapy (Felsberg 2009). In all other studies, we had low concerns or were uncertain. Full details of the applicability assessments and their justifications are available in Appendix 7.

Findings: comparisons of different methods

We provide all of the 160 HRs we extracted in Appendix 8. In all cases, the estimated HR was above 1, indicating higher hazard of death in people with an unmethylated MGMT promoter (or MGMT protein expression on IHC). In the vast majority of cases, the lower limit of a 95% CI for the HR was above 1, confirming the prognostic value of MGMT promoter methylation status. When examining these results and the forest plots that follow, we emphasise that

comparisons should only be made of different methods *within* studies. HRs should not be compared across studies because there were many (more substantial) differences between these results than the choice of technique, tumour sample, CpG islands or cut-off thresholds.

Comparison of methylation-specific polymerase chain reaction versus immunohistochemistry

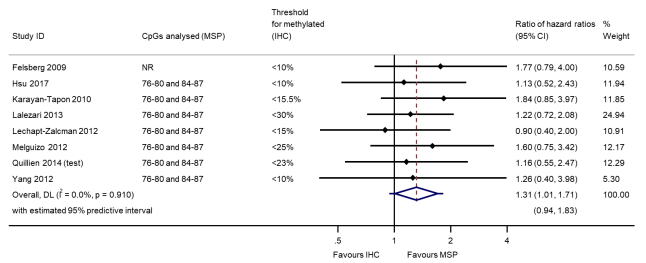
We illustrate results from eight studies that included both MSP and IHC in Figure 4. Seven reported they had targeted CpG sites 76 to 80 and 84 to 87 in MSP (the other did not report their CpG sites). Risk of bias was assessed to be low (or unclear) in all of these results except in one study where the cut-off threshold for interpretation of IHC for the MGMT protein was based on its performance (Quillien 2014 (test)), which we assessed to be at high risk of bias. There was a tendency for MSP to produce larger HRs than IHC, suggesting that MSP provides a more discriminating predictor of time to death than IHC for MGMT protein. RHRs providing direct within-study comparisons of MSP versus IHC for these studies are presented in Figure 5. A meta-analysis of these gave a summary RHR of 1.31 (95% CI 1.01 to 1.71), providing some statistical support for the observation that MSP is more prognostic for overall survival than IHC for MGMT protein. There was no evidence of heterogeneity between the studies (estimated between-study variance = 0; I^2 = 0%), and a 95% prediction interval from 0.94 to 1.83 was therefore barely any wider than the CI.

Figure 4. Hazard ratios from studies comparing MSP with IHC. CI: confidence interval; CpG: 5'-cytosine-phosphateguanine-3'; FFPE: formalin-fixed paraffin embedded; HR: hazard ratio; IHC: immunohistochemistry; MSP: methylation-specific polymerase chain reaction; N/A: not applicable; NR: not reported; PCR: polymerase chain reaction; PF: prognostic factor; RoB: risk of bias; sel. rep.: selective reporting.

Study ID and Technique Sample	CpGs analysed type (PCR-based tests)	Threshold for methylated			Hazard ratio (95% CI)	Sample size	D4: PF	D5: attrition	D7: sel. rep.
Felsberg 2009 IHC FFPE MSP Frozen	N/A (14 FFPE) NR	<10% NR			1.26 (0.70, 2.25) 2.23 (1.29, 3.84)	61 66	Low RoB Low RoB	Low RoB Low RoB	Low RoB Low RoB
Hsu 2017 IHC FFPE MSP FFPE	N/A 76-80 and 84-87	<10% NR			2.12 (1.32, 3.42) 2.39 (1.42, 4.02)	121 121	Low RoB Low RoB	Low RoB Low RoB	Low RoB Low RoB
Karayan-Tapon 2010 IHC FFPE MSP Frozen	N/A 76-80 and 84-87	<15.5% NR		└ ╋ <u></u>	1.26 (0.77, 2.06) 2.32 (1.39, 3.87)	78 81	Low RoB Low RoB	Low RoB Low RoB	Low RoB Low RoB
Lalezari 2013 IHC FFPE MSP FFPE	N/A 76-80 and 84-87	<30% NR		→	1.74 (1.39, 2.16) 2.13 (1.67, 2.78)	355 402	Low RoB Low RoB	Unclear Low RoB	Low RoB Low RoB
Lechapt-Zalcman 20 IHC FFPE MSP FFPE	12 N/A 76-80 and 84-87	<15% NR		+	1.99 (1.15, 3.42) 1.78 (1.03, 3.09)	106 110	Low RoB Low RoB	Low RoB Low RoB	Low RoB Low RoB
Melguizo 2012 IHC FFPE MSP NR	N/A 76-80 and 84-87	<25% NR		• <u> </u> • •	1.11 (0.69, 1.77) 1.77 (1.06, 2.95)	76 76	Low RoB Low RoB	Low RoB Low RoB	Low RoB Low RoB
Quillien 2014 (test) IHC FFPE MSP Frozen	N/A 76-80 and 84-87	<23% NR			2.33 (1.44, 3.74) 2.70 (1.65, 4.43)	99 99	High RoB Low RoB	Low RoB Low RoB	Low RoB Low RoB
Yang 2012 IHC FFPE MSP FFPE	N/A 76-80 and 84-87	<10%		• •	1.07 (0.35, 3.31) 1.35 (0.44, 4.16)	18 18	Low RoB Low RoB	Low RoB Low RoB	Unclear Unclear
			.5	I 1 2	4				

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)

Figure 5. Meta-analysis of ratios of hazard ratios (RHR) providing within-study comparisons of MSP and IHC (an RHR greater than 1 indicates that MSP is more prognostic than IHC). CpG: 5'-cytosine-phosphate-guanine-3'; CI: confidence interval; DL: DerSimonian and Laird estimation; IHC: immunohistochemistry; MSP: methylation-specific polymerase chain reaction.



NOTE: Weights are from random-effects model

Technical notes: in the four data sets from the whole review for which we had IPD, the correlations between results on pairs of test results for the same people (categorised as methylated versus unmethylated) were 0.75 (MSP versus PSQ; McDonald 2013), 0.88 (MSP versus bead array; Bady 2012 (M-GBM)), 0.65 (IHC versus qMSP; Hsu 2015) and -0.03 (MSP versus IHC; Yang 2012). Based on these observations, in our statistical comparison of HRs between methods within the same study, we used a value of 0.7 for the correlation, and repeated the analysis using correlations of 0.5 and 0.9 as our sensitivity analysis.

Sensitivity analysis: when we assumed a correlation of 0.5, the RHR was 1.31 (95% CI 0.93 to 1.85); when we assumed a correlation of 0.9, the RHR was 1.32 (95% CI 1.12 to 1.56). There was a small amount of heterogeneity in the latter analysis (between-study

variance = 0.008) due to the narrowing of CIs for the RHRs in individual studies.

Comparison of pyrosequencing versus immunohistochemistry

We illustrate results from five studies that included both PSQ and IHC for MGMT protein in Figure 6. Risks of bias were variable, with seven of the 10 results being at low or unclear risk of bias in all domains. In one study (in which cut-off thresholds were derived based on performance) both results were assessed to be at high risk of bias (Quillien 2014 (test)), and in another study we rated the PSQ result to be at high risk of bias in the attrition domain because it omitted more than 25% of the sample without explanation (Lalezari 2013). In all the studies, IHC had been performed using FFPE samples, but in three PSQ was performed using frozen samples.

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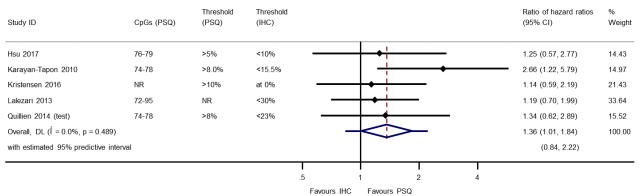
Figure 6. Hazard ratios from studies comparing PSQ with IHC. CI: confidence interval; CpG: 5'-cytosine-phosphateguanine-3'; FFPE: formalin-fixed paraffin embedded; HR: hazard ratio; IHC: immunohistochemistry; N/A: not applicable; NR: not reported; PCR: polymerase chain reaction; PF: prognostic factor; PSQ: pyrosequencing; RoB: risk of bias; sel. rep.: selective reporting

Study ID an Technique	d Sample type	CpGs analysed (PCR-based tests)	Threshold for methylated				Hazard ratio (95% CI)	Sample size	D4: PF	D5: attrition	D7: sel. rep.
Hsu 2017 IHC PSQ	FFPE FFPE	N/A 76-79	<10% >5%		* *		2.12 (1.32, 3.42) 2.66 (1.49, 4.76)	121 121	Low RoB Low RoB	Low RoB Low RoB	Low RoB Low RoB
Karayan-Ta IHC PSQ	pon 2010 FFPE Frozen	N/A 74-78	<15.5% >8.0%	•			1.26 (0.77, 2.06) 3.35 (1.95, 5.73)	78 79	Low RoB Low RoB	Low RoB Low RoB	Low RoB Low RoB
Kristensen IHC PSQ	2016 FFPE Frozen	N/A NR	at 0% ≻10%				1.58 (1.12, 2.22) 1.80 (1.23, 2.62)	148 132	Unclear Low RoB	Low RoB Unclear	Low RoB Low RoB
Lalezari 20 [.] IHC PSQ	I3 FFPE FFPE	N/A 72-95	<30% NR	│ <u>→</u>	→		1.74 (1.39, 2.16) 2.06 (1.62, 2.62)	355 312	Low RoB Low RoB	Unclear High RoB	Low RoB Low RoB
Quillien 201 IHC PSQ	4 (test) FFPE Frozen	N/A 74-78	<23% >8%		++		2.33 (1.44, 3.74) 3.13 (1.86, 5.25)	99 99	High RoB High RoB	Low RoB Low RoB	Low RoB Low RoB
			I .5	1	2	I 4					

RHRs providing direct within-study comparisons of PSQ versus IHC for these studies are presented in Figure 7. There was evidence of superiority of PSQ over IHC (RHR 1.36, 95% CI 1.01 to 1.84). There was no evidence of statistical heterogeneity (between-study

variance = 0; $I^2 = 0\%$; the wide 95% prediction interval from 0.84 to 2.22 represents uncertainty in estimation of the between-study variance).

Figure 7. Meta-analysis of ratios of hazard ratios (RHR) providing within-study comparisons of PSQ and IHC (an RHR greater than 1 indicates that PSQ is more prognostic than IHC). CI: confidence interval; CpG: 5'-cytosine-phosphate-guanine-3'; DL: DerSimonian and Laird estimation; IHC: immunohistochemistry; NR: not reported; PSQ: pyrosequencing.



NOTE: Weights are from random-effects model

Technical notes: four studies had used more than three variants of PSQ, and all of these had examined CpG sites 74 to 78 with a cutoff of 8%, applied to frozen tumour samples (Havik 2012; Karayan-Tapon 2010; Quillien 2014 (test); Quillien 2016). This combination had also been used in two other studies comparing PSQ with MSP (Bady 2012 (E-GBM); Kim 2016). Therefore, we included only this combination from these four studies in the forest plot. (Detailed comparison of PSQ variants are covered below).

Sensitivity analysis: when we assumed a correlation of 0.5, the RHR was 1.36 (95% CI 0.92 to 2.01); when we assumed a correlation of 0.9, the RHR was 1.41 (95% CI 1.06 to 1.87). Between-study variance was estimated to be 0.06 in the latter analysis.

Comparison of methylation-specific polymerase chain reaction versus pyrosequencing

We illustrate results from nine studies that included both MSP and PSQ in Figure 8. Three of the 22 results in this plot were considered to be at high risk of bias: for Havik 2012 and Quillien 2014 (test) due to a data-based selection of a threshold for PSQ, and for Lalezari 2013 due to missing data for PSQ. RHRs providing direct within-study comparisons of MSP versus PSQ for these studies are presented in Figure 9. While there was a consistent pattern that PSQ seemed to be a slightly better predictor than MSP, there was not strong statistical evidence to confirm this (RHR 1.14, 95% CI 0.87 to 1.48). There was no evidence of statistical heterogeneity (between-

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study variance = 0; l^2 = 0%), with a 95% prediction interval for the HRH ranging from 0.83 to 1.56. No pattern could be discerned in a

funnel plot of these results (Figure 10); tests for asymmetry were not undertaken.

Figure 8. Hazard ratios from studies comparing PSQ with MSP. CI: confidence interval; CpG: 5'-cytosine-phosphateguanine-3'; FFPE: formalin-fixed paraffin embedded; HR: hazard ratio; MSP: methylation-specific polymerase chain reaction; N/A: not applicable; NR: not reported; PCR: polymerase chain reaction; PF: prognostic factor; PSQ: pyrosequencing; RoB: risk of bias; sel. rep.: selective reporting

Study ID and Technique	Sample type	CpGs analysed (PCR-based tests)	Threshold for methylated		Hazard ratio (95% CI)	Sample size	D4: PF	D5: attrition	D7: sel. rep.
Barbagallo 20 MSP MSP PSQ PSQ	14 FFPE FFPE FFPE FFPE	76-80 and 84-87 76-80 and 84-87 NR NR	Incl. weakly Excl. weakly >9% >25%		3.68 (1.66, 8.18) 1.90 (0.72, 4.99) 3.73 (1.68, 8.28) 1.99 (0.92, 4.32)	37 37 37 37	Low RoB Low RoB Low RoB Low RoB	Low RoB Low RoB Low RoB Low RoB	Low RoB Low RoB Low RoB Low RoB
Havik 2012 MSP PSQ	Frozen Frozen	76-80 and 84-87 74-78	NR >8%		2.02 (1.08, 3.77) 2.33 (1.19, 4.57)	48 48	Low RoB High RoB	Low RoB Low RoB	Low RoB Low RoB
Hsu 2017 MSP PSQ	FFPE FFPE	76-80 and 84-87 76-79	NR >5%		2.39 (1.42, 4.02) 2.66 (1.49, 4.76)	121 121	Low RoB Low RoB	Low RoB Low RoB	Low RoB Low RoB
Karayan-Tapo MSP PSQ	n 2010 Frozen Frozen	76-80 and 84-87 74-78	NR >8.0%	— — —	2.32 (1.39, 3.87) 3.35 (1.95, 5.73)	81 79	Low RoB Low RoB	Low RoB Low RoB	Low RoB Low RoB
Kim 2016 MSP PSQ	FFPE FFPE	76-80 and 84-87 74-78	NR >9%		7.66 (2.82, 20.81) 7.66 (2.82, 20.81)		Low RoB High RoB	Low RoB Low RoB	Low RoB Low RoB
Lalezari 2013 MSP PSQ	FFPE FFPE	76-80 and 84-87 72-95	NR NR		2.13 (1.67, 2.78) 2.06 (1.62, 2.62)	402 312	Low RoB Low RoB	Low RoB High RoB	Low RoB Low RoB
Lattanzio 2015 MSP MSP PSQ PSQ	FFPE Frozen FFPE Frozen	76-80 and 84-87 76-80 and 84-87 72-80 72-80	NR NR ?9% ?9%		1.45 (0.76, 2.76) 2.27 (1.21, 4.26) 2.09 (1.09, 3.99) 2.25 (1.19, 4.25)	40 45 45 45	Low RoB Low RoB Low RoB Low RoB	Unclear Low RoB Low RoB Low RoB	Low RoB Low RoB Low RoB Low RoB
McDonald 201 MSP PSQ	3 FFPE FFPE	76-80 74-78	NR >8%		1.64 (0.95, 2.83) 1.96 (1.16, 3.33)	71 76	Low RoB Low RoB	Unclear Low RoB	Low RoB Low RoB
Quillien 2014 MSP PSQ	(test) Frozen Frozen	76-80 and 84-87 74-78	NR >8%		2.70 (1.65, 4.43) 3.13 (1.86, 5.25)	99 99	Low RoB High RoB	Low RoB Low RoB	Low RoB Low RoB
			I .5	I I 1 2 4					

Figure 9. Meta-analysis of ratios of hazard ratios (RHR) providing within-study comparisons of PSQ and MSP (an RHR greater than 1 indicates that PSQ is more prognostic than MSP). CI: confidence interval; CpG: 5'-cytosine-phosphate-guanine-3'; HR: hazard ratio; MSP: methylation-specific polymerase chain reaction; NR: not reported; PSQ: pyrosequencing.

Study ID	CpGs (PSQ)	CpGs (MSP)	Threshold (PSQ)	Threshold (MSP)					Ratio of hazard ratios (95% CI)	% Weight
Barbagallo 2014	NR	76-80 and 84-87	>9%	Incl. weakly					1.01 (0.38, 2.67)	7.41
Havik 2012	74-78	76-80 and 84-87	>8%	NR					1.16 (0.48, 2.78)	9.09
Hsu 2017	76-79	76-80 and 84-87	>5%	NR	-				1.11 (0.50, 2.49)	10.68
Karayan-Tapon 2010	74-78	76-80 and 84-87	>8.0%	NR		-+	•		1.44 (0.66, 3.17)	11.24
Kim 2016	74-78	76-80 and 84-87	>9%	NR -		∔ ¦			1.00 (0.34, 2.96)	5.91
Lalezari 2013	72-95	76-80 and 84-87	NR	NR		•			0.97 (0.56, 1.66)	23.86
Lattanzio 2015	72-80	76-80 and 84-87	?9%	NR		;-	•		1.44 (0.60, 3.45)	9.12
McDonald 2013	74-78	76-80	>8%	NR		-++			1.20 (0.54, 2.64)	11.04
Quillien 2014 (test)	74-78	76-80 and 84-87	>8%	NR		-++-			1.16 (0.53, 2.50)	11.65
Overall, DL ² (# 0.0%, p = 0.9	997)					$ \leftarrow $	>		1.14 (0.87, 1.48)	100.00
with estimated 95% predict	tive interval								(0.83, 1.56)	
					1			I	I	
					.5	1		2	4	
					Favours MSP		Favours PS	Q		

NOTE: Weights are from random-effects model

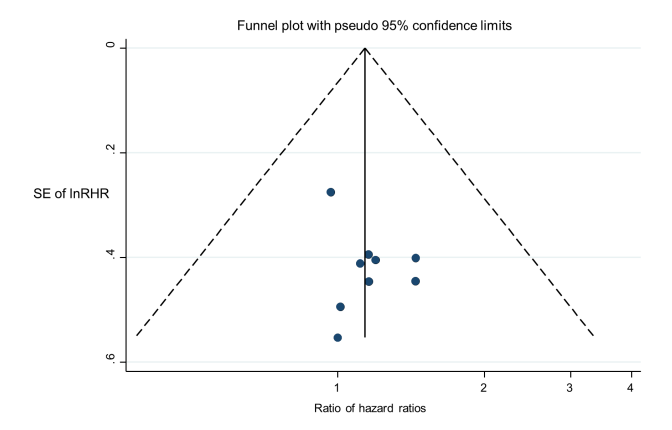
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Figure 10. Funnel plot of ratios of hazard ratios (RHR) providing within-study comparisons of PSQ and MSP (an RHR greater than 1 indicates that PSQ is more prognostic than MSP). MSP: methylation-specific polymerase chain reaction; PSQ: pyrosequencing; SE: standard error.



Technical notes: we restricted these forest plots (and funnel plot) to PSQ results for CpG sites 74 to 78 with a cut-off of 8% and frozen tumour samples for four studies (Havik 2012; Karayan-Tapon 2010; Quillien 2014 (test); Quillien 2016). To select a single HR for each technique in each study to estimate RHRs, we applied two further rules: we selected FFPE over frozen (as it was used more often across studies), and a cut-off for PSQ of less than 9% over less than 25% (for consistency with other studies).

Sensitivity analysis: when we assumed a correlation of 0.5, the RHR was 1.14 (95% CI 0.81 to 1.60); when we assumed a correlation of 0.9, the RHR was 1.14 (95% CI 0.98 to 1.33). Heterogeneity was estimated to be 0 in these analyses. Thus, assuming HRs for MSP and PSQ were highly correlated (0.9), the CI narrowed sufficiently to be suggestive of a difference in favour of PSQ.

Quantitative methylation-specific polymerase chain reaction

Figure 11 shows HRs for studies comparing different methods for qMSP, or comparing qMSP against either MSP or PSQ. The qMSP methods most commonly targeted CpG sites 76 to 80 and 84 to 87. Quillien 2016 and Yoshioka 2018 both compared SQ-MSP, using frozen tissue samples, targeting CpGs 76 to 80 and 84 to 87 using different thresholds. Both observed that HRs were higher the lower the cut-off point. Quillien 2016 observed the opposite when they looked also at FFPE samples. Across the studies, there was no indication that qMSP methods were superior to PSQ. The one study that looked at MethyLight-MSP did not find it very prognostic (Quillien 2014 (test)).

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Figure 11. Hazard ratios from studies comparing different methods for qMSP, or comparing qMSP against either MSP or PSQ. CI: confidence interval; CpG: 5'-cytosine-phosphate-guanine-3'; FFPE: formalin-fixed paraffin embedded; FSQ-MS-PCR: fluorescent semi-quantitative methylation-specific polymerase chain reaction; HR: hazard ratio; MSP: methylation-specific polymerase chain reaction; NR: not reported; PCR: polymerase chain reaction; PF: prognostic factor; PSQ: pyrosequencing; qMSP: quantitative methylation-specific polymerase chain reaction; RoB: risk of bias; sel. rep.: selective reporting; SQ-MSP: semi quantitative methylation-specific polymerase chain reaction.

Heak 2012 MSP Frozen 74-50 and 84-87 NR PSO Frozen 74-78 >8% NR 2.02 (1 08, 3.77) 48 Low RoB	Study ID and Technique	Sample type	CpGs analysed (PCR-based tests)	Threshold for methylated				Hazard ratio (95% CI)	Sample size	D4: PF	D5: attrition	D7: sel. rep.
PSQ GMSP Frozen Frozen 74-78 71-78 and 75-86 >8% GMSP Frozen Frozen 71-78 and 75-86 NR SPG GMSP FFPE Frozen 76-80 and 84-87 NR SPS FFPE GMSP 77-80 and 84-87 NR SPS FFPE Frozen 77-80 and 84-87 NR SSD-MSP FFPE Frozen 77-80 and 84-87 NR SGMSP FFPE 77-80 and 84-87 NR SGMSP FFPE 77-80 and 84-87 NR SGMSP Frozen 76-80 and 84-87 NR SGMSPCR Frozen 76-80 and 84-87 NR SGMSPCR Frozen 76-80 and 84-87 >0.0% SGMSPCR Frozen 76-80 and 84-87 >15% SGMSPCR Frozen 76-80 and 84-87 >15% SGMSPCR Frozen 76-80 and 84-87 >15% SGMSPCR Frozen 76-80 and 84-87 >15% 228 (1.70, 4.21) 106 High RoB Low RoB Low RoB Low RoB SQMSPCR Frozen 76-80 and 84-87 >15% 20 270 (1.65, 4.43) <th< td=""><td></td><td></td><td></td><td></td><td></td><td>•</td><td></td><td></td><td></td><td></td><td></td><td></td></th<>						•						
gMSP Frozen 71-73 and 75-86 NR gMSP Frozen 71-88 >>0% 172 (0.91, 3.22) 48 Low RoB Low RoB Low RoB MSP FFPE 76-80 and 84-87 NR 239 (1.42, 4.02) 121 Low RoB Low Ro						•						
dKSP Frozen 71-86 >0% 166 (0.97, 2.83) 58 Low RoB Low RoB Low RoB Hsu 2017 MSP FFPE 76-79 >5% 100 239 (142, 4.02) 121 Low RoB Low RoB Low RoB gMSP FFPE 77-90 and 84-87 >0.04% 239 (142, 4.02) 121 Low RoB Low RoB Low RoB gMSP FFPE 77-90 and 84-87 >0.04% 233 (145, 5.73) 79 Low RoB Low RoB Low RoB SG-MSPCR Frozen 74-80 and 84-87 >0.04% >0.05% 232 (1.39, 3.87) 81 Low RoB Low RoB Low RoB SG-MSPCR Frozen or FFPE 76-80 and 84-87 >55% 25% 225 (1.31, 3.47) 106 High RoB Low RoB Low RoB SG-MSPCR Frozen or FFPE 76-80 and 84-87 >15% >0 25% 106 High RoB Low RoB					-	•						
Hsu 2017 MSP FFPE 76-80 and 84-87 NR MSP FFPE 77-80 and 84-87 >0.04% MSP Frozen 76-80 and 84-87 >0.04% SQ-MSP Frozen 76-80 and 84-87 >35 SQ-MSP Frozen 76-80 and 84-87 >35 SQ-MSP CR Frozen 76-80 and 84-87 >56% SQ-MSP FCR Frozen 76-80 and 84-87 >6% SQ-MSP FCR Frozen 76-80 and 84-87 >6% SQ-MSP FCR Frozen 76-80 and 84-87 >6% SQ-MSP FCR Frozen 76-80 and 84-87 >8% SQ-MSP Frozen 76-80 and 84-87 >8% >6% SQ-MSP Frozen 76-80 and 84-87<												
MSP FFPE 76-80 and 84-87 NR gMSP FFPE 77-80 and 84-87 > 00 4% gMSP FFPE 77-80 and 84-87 > 01 0% gMSP FFPE 77-80 and 84-87 > 01 0% gMSP FFPE 77-80 and 84-87 > 01 % gMSP FFPE 77-80 and 84-87 > 01 % gMSP FFPE 77-80 and 84-87 > 01 % gMSP FFPE 77-80 and 84-87 > 0.1% gMSP Frozen 76-80 and 84-87 > 80 % gSUAMSP Frozen or FFPE 76-80 and 84-87 > 55% SQMSP-C Frozen or FFPE 76-80 and 84-87 > 15% SQMSP-C Frozen or FFPE 76-80 and 84-87 > 15% SQMSP-C Frozen or FFPE 76-80 and 84-87 > 15% SQMSP-C Frozen or FFPE 76-80 and 84-87 > 15% SQMSP-C Frozen or FFPE 76-80 and 84-87 > 15% SQMSP-C Frozen or FFPE 76-80 and 84-87 > 15% SQMSP-C Frozen or FFPE 76-80 and 84-87 > 15% SQ-MSP	qMSP	Frozen	/1-86	>0%				1.66 (0.97, 2.83)	58	Low RoB	Low RoB	Low RoB
PSQ FFPE 76-79 >5% gMSP FFPE 77-80 and 84-87 >0.04% gMSP FFPE 77-80 and 84-87 >0.04% 2.75 (15.15, 04) (121 Low RoB Low RoB Low RoB Low RoB Low RoB Karayan-Tapon 2010 MSP Frozen 76-80 and 84-87 >0.1% 2.32 (139, 3.87) 81 Low RoB Low RoB Low RoB Low RoB Low RoB SQ-MSP Frozen 76-80 and 84-87 >8.0% 2.32 (139, 3.87) 81 Low RoB Low												
gMSP FFPE 77-80 and 84-87 >0.04% 2.75 (15.15.0.4) 121 Low RoB Low RoB Low RoB MSP FFPE 77-80 and 84-87 >0.1% 2.33 (1.85, 4.33) 118 Low RoB						•						
qMSP FFPE 77-80 and 84-87 >0.1% 2.83 (1.85, 4.33) 118 Low RoB Low RoB Low RoB Karayan-Tapon 2010 MSP Frozen 76-80 and 84-87 NR >8.0% 2.32 (1.39, 3.87) 81 Low RoB												
Karayan-Tapon 2010 NR Pozen 76-80 and 84-87 NR 23.25 (1.39, 3.87) 81 Low RoB L												
MSP Frozen 76-80 and 84-87 NR SQ-MSP Frozen 76-80 and 84-87 >356 SQ-MSP Frozen 76-80 and 84-87 >356 SQ-MSP Frozen 76-80 and 84-87 >356 FSQ-MS-PCR Frozen or FFPE 76-80 and 84-87 >15% SQ-MSP Frozen or FFPE 76-80 and 84-87 >15% SQ-MSP Frozen or FFPE 76-80 and 84-87 >15% Methylight-MSP Frozen or FFPE 76-80 and 84-87 >15% PSQ Frozen or FFPE 76-80 and 84-87 >8% Methylight-MSP Frozen or FFPE 76-80 and 84-87 >8% PSQ FFPE 74-78 >8% PSQ FFPE 74-78 >8% SQ-MSP Frozen or 74-78 >8% >8% 270 (16.5, 4.43) 99 Low RoB <	qMSP	FFPE	77-80 and 84-87	>0.1%	-			2.83 (1.85, 4.33)	118	Low RoB	Low RoB	Low RoB
PSQ Frozen 74-78 >80% SQ-MSP Frozen 76-80 and 84-87 >35 Nguyen 2015 FSQ-MS-PCR Frozen or FFPE 76-80 and 84-87 >15% FSQ-MS-PCR Frozen or FFPE 76-80 and 84-87 >60% 2.68 (1.70, 4.21) 106 High RoB Low RoB Low RoB Low RoB Quillen 2014 (test) Frozen or FFPE 76-80 and 84-87 >60% 2.70 (1.65, 4.43) 99 Low RoB												
SQ-MSP Frozen 76-80 and 84-87 >35 2.75 (1.66, 4.53) 81 Low RoB Low RoB Low RoB Nguyen 2015 FSQ-MS-PCR Frozen or FFPE 76-80 and 84-87 >15% 2.68 (1.70, 4.21) 106 High RoB Low RoB						-						
Nguyan 2015 FSQ-MS-PCR Frozen or FFPE Frozen or FFPE 76-80 and 84-87 76-80 and 84-87 >15% >60% Quillien 2014 (test) MSP msp Frozen or FFPE 76-80 and 84-87 76-80 and 84-87 NR >0% Quillien 2014 (test) MSP msp Frozen or FFPE 76-80 and 84-87 76-80 and 84-87 NR >0 SQ Frozen or FFPE 76-80 and 84-87 76-80 and 84-87 NR >0 270 (165, 443) 99 167 (100, 2.77) Low RoB 167 (100, 2.77) Low RoB 167 (100, 2.77) Low RoB 167 (100, 2.77) Low RoB 169 (10, 2.77) Low RoB 100 RoB Low RoB Low RoB Quillien 2016 PSQ Frozen or 74-78 Frozen 74-78 >8% ++++++++++++++++++++++++++++++++++++							•					
FSQ-MS-PCR Frozen or FFPE 76-80 and 84-87 >15% Guillien 2014 (test) MSP Frozen or FFPE 76-80 and 84-87 >15% MSP Frozen or FFPE 76-80 and 84-87 NR MSP Frozen or FFPE 76-80 and 84-87 NR MSP Frozen or FFPE 76-80 and 84-87 NR PSQ Frozen or FFPE 76-80 and 84-87 NR PSQ Frozen or FFPE 74-78 >8% Quillien 2016 PSQ Frozen or 74-78 >8% PSQ FFPE 76-80 and 84-87 >12% 4.00 (2.30, 6.97) 121 Low RoB Low RoB Low RoB Quillien 2016 PSQ Frozen or 74-78 >8% >3.33 (2.06, 5.40) 118 Unclear Low RoB Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >13% >23% 23% 23% 23% 23% 23% 24% 23% 24% 23% 24% 23% 23% 24% 23% 24% 23% 24% 26% 277 (1.85, 7.38) 118 Unclear Low RoB <	SQ-MSP	Frozen	76-80 and 84-87	>35		+		2.75 (1.66, 4.53)	81	Low RoB	Low RoB	Low RoB
FSQ-MS-PCR Frozen or FFPE 76-80 and 84-87 >60% 2.25 (1.31, 3.87) 106 High RoB Low RoB Low RoB Quillien 2014 (test) MSP Frozen 76-80 and 84-87 NR 2.70 (1.85, 4.43) 99 Low RoB Low RoB Low RoB Low RoB PSQ Frozen 74-78 >8% 167 (1.00, 2.77) 99 High RoB Low RoB Low RoB Low RoB Quillien 2016 PSQ Frozen 74-78 >8% 4.00 (2.30, 6.97) 121 Low RoB Low RoB Low RoB SQ-MSP FFPE 76-80 and 84-87 >13% >8% 3.33 (2.06, 5.40) 118 Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >13% >3% 2.33 (2.06, 5.40) 118 Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >13% 23% 2.33 (2.06, 5.40) 118 Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% 2.33 (2.06, 5.40) 118 Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23%	Nguyen 2015											
Quillien 2014 (lest) MSP Frozen 76-80 and 84-87 NR MethylLight-MSP Frozen 75-86 >0 PSQ Frozen 74-78 >8% Quillien 2016 PSQ Frozen 74-78 >8% PSQ Frozen 74-78 >8% Quillien 2016 PSQ Frozen 74-78 >8% PSQ Frozen 74-78 >8% SQ-MSP FFPE 76-80 and 84-87 >13% SQ-MSP FFPE 76-80 and 84-87 >13% SQ-MSP FFPE 76-80 and 84-87 >23% SQ-MSP Frozen 76-80 and 84-87	FSQ-MS-PCR	Frozen or FFPE	76-80 and 84-87	>15%		•		2.68 (1.70, 4.21)	106	High RoB	Low RoB	Low RoB
MSP Frozen 76-80 and 84-87 NR MethylLight-MSP Frozen 75-86 >0 PSQ Frozen 74-78 >8% Quillien 2016 PSQ Frozen 74-78 >8% PSQ Frozen 74-78 >8% 4.00 (2.30, 6.97) 121 Low RoB Low RoB Low RoB Quillien 2016 PSQ Frozen 74-78 >8% 3.13 (1.86, 5.25) 99 High RoB Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >13% 3.33 (2.06, 5.40) 118 Unclear Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >13% 3.33 (2.06, 5.40) 118 Unclear Unclear Low RoB SQ-MSP Frpzen 76-80 and 84-87 >23% 2.66 (1.79, 4.57) 18 High RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% 2.17 (1.38, 3.44) 108 High RoB Unclear Low RoB SQ-MSP <t< td=""><td>FSQ-MS-PCR</td><td>Frozen or FFPE</td><td>76-80 and 84-87</td><td>>60%</td><td></td><td>•</td><td></td><td>2.25 (1.31, 3.87)</td><td>106</td><td>High RoB</td><td>Low RoB</td><td>Low RoB</td></t<>	FSQ-MS-PCR	Frozen or FFPE	76-80 and 84-87	>60%		•		2.25 (1.31, 3.87)	106	High RoB	Low RoB	Low RoB
MethylLight-MSP Frozen 75-86 >0 PSQ Frozen 74-78 >8% Quillen 2016 PSQ FFPE 74-78 >8% PSQ FFPE 74-78 >8% 4.00 (2:30, 6:97) 121 Low RoB Low RoB Low RoB Quillen 2016 PSQ FFPE 74-78 >8% 3.57 (2:14, 5:95) 112 Low RoB Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >13% 3.33 (2:06, 5:40) 118 Unclear Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >23% 3.33 (2:06, 5:40) 118 Unclear Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% >3.33 (2:06, 5:40) 118 Unclear Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% >23% 2.17 (1:38, 3:44) 108 High RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% 2.16 (1:73, 4:57) 80 Low RoB Unclear Low RoB	Quillien 2014 (te	st)										
PSQ Frozen 74-78 >8% Quillien 2016 PSQ FFPE 74-78 >8% PSQ Frozen 74-78 >8% SQ_MSP FFPE 74-78 >8% SQ_MSP FFPE 76-80 and 84-87 >12% Low RoB Unclear Low RoB SQ_MSP FFPE 76-80 and 84-87 >12% 33 (2 06, 5.40) 118 Unclear Unclear Low RoB SQ_MSP FFPE 76-80 and 84-87 >13% 2.36 (1.79, 4.57) 108 High RoB Unclear Low RoB SQ_MSP Frozen 76-80 and 84-87 >23% 2.86 (1.79, 4.57) 108 High RoB Unclear Low RoB SQ_MSP Frozen 76-80 and 84-87 >23% 2.17 (1.38, 3.44) 108 High RoB Unclear Low RoB SQ_MSP Frozen 76-80 and 84-87 >4 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ_MSP Frozen 76-80 and 84-87 >4 2.72 (MSP	Frozen	76-80 and 84-87	NR				2.70 (1.65, 4.43)	99	Low RoB	Low RoB	Low RoB
Quillien 2016 PSQ FFPE 74-78 >8% PSQ Frozen 74-78 >8% SQ-MSP FFPE 76-80 and 84-87 >12% Low RoB Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >13% 333 (2 06, 5.40) 118 Unclear Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >23% 333 (2 06, 5.40) 118 Unclear Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >23% 2.86 (1.79, 4.57) 108 High RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% 2.17 (1.33, 3.44) 108 High RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >2 2.17 (1.33, 3.44) 108 High RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >4 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87	MethylLight-MSP	Frozen	75-86	>0				1.67 (1.00, 2.77)	99	High RoB	Low RoB	Low RoB
PSQ FFPE 74-78 >8% PSQ Frpzen 74-78 >8% PSQ Frozen 74-78 >8% SQ-MSP FFPE 76-80 and 84-87 >12% 4.00 (2.30, 6.97) 121 Low RoB Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >12% 3.33 (2.06, 5.40) 118 Unclear Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >23% 23% 4.17 (2.35, 7.38) 118 Unclear Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% 23% 4.17 (2.35, 7.38) 118 Unclear Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% 23% 4.17 (2.35, 7.38) 118 Unclear Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% 23% 2.17 (1.38, 3.44) 108 High RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >4 2 2 2.17 (1.38, 3.44) 80 Low RoB	PSQ	Frozen	74-78	>8%	-	•		3.13 (1.86, 5.25)	99	High RoB	Low RoB	Low RoB
PSQ Frozen 74-78 >8% SQ-MSP FFPE 76-80 and 84-87 >12% SQ-MSP FFPE 76-80 and 84-87 >13% SQ-MSP FFPE 76-80 and 84-87 >13% SQ-MSP FFPE 76-80 and 84-87 >23% SQ-MSP FFPE 76-80 and 84-87 >23% SQ-MSP Frozen 76-80 and 84-87 >23% SQ-MSP Frozen 76-80 and 84-87 >23% SQ-MSP Frozen 76-80 and 84-87 >23% Yoshioka 2018 2018 217 (1.38, 3.44) 108 High RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >0 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >0 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >0 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >6 2.72 (1.28, 5.74) 80	Quillien 2016											
SQ-MSP FFPE 76-80 and 84-87 >12% SQ-MSP FFPE 76-80 and 84-87 >13% SQ-MSP FFPE 76-80 and 84-87 >333 (2.06, 5.40) 118 Unclear Unclear Low RoB SQ-MSP FFPE 76-80 and 84-87 >23%	PSQ	FFPE	74-78	>8%				4.00 (2.30, 6.97)	121	Low RoB	Unclear	Low RoB
SQ-MSP FFPE 76-80 and 84-87 >13% SQ-MSP FFPE 76-80 and 84-87 >23% SQ-MSP FFPE 76-80 and 84-87 >23% SQ-MSP Frozen 76-80 and 84-87 >13% SQ-MSP Frozen 76-80 and 84-87 >23% Yoshioka 2018 2018 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB Yoshioka 2018 SQ-MSP Frozen 76-80 and 84-87 >2 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >4 76-80 and 84-87 >4 76-80 and 84-87 >2 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >4 4 76-80 and 84-87 >4 4 76-80 and 84-87 >6 1.85 (1.07, 3.18) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 <td< td=""><td>PSQ</td><td>Frozen</td><td>74-78</td><td>>8%</td><td></td><td></td><td>•</td><td>3.57 (2.14, 5.95)</td><td>112</td><td>Low RoB</td><td>Unclear</td><td>Low RoB</td></td<>	PSQ	Frozen	74-78	>8%			•	3.57 (2.14, 5.95)	112	Low RoB	Unclear	Low RoB
SQ-MSP FFPE 76-80 and 84-87 >23% SQ-MSP Frozen 76-80 and 84-87 >1% +17 (2.35, 7.38) 118 High RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% 2.17 (1.38, 3.44) 108 High RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >23% 2.17 (1.38, 3.44) 108 High RoB Unclear Low RoB Yoshioka 2018 SQ-MSP Frozen 76-80 and 84-87 >0 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >0 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >4 1.85 (1.07, 3.18) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >6 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >8 1.83 (1.10,	SQ-MSP	FFPE	76-80 and 84-87	>12%			•	3.33 (2.06, 5.40)	118	Unclear	Unclear	Low RoB
SQ-MSP Frozen 76-80 and 84-87 >13% SQ-MSP Frozen 76-80 and 84-87 >23% Yoshioka 2018 2.17 (1.38, 3.44) 108 High RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >2 2.17 (1.38, 3.44) 108 High RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >2 2.18 (1.20, 3.97) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >2 2.18 (1.07, 3.18) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >4 1.85 (1.07, 3.18) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >4 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >6 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >6	SQ-MSP	FFPE	76-80 and 84-87	>13%			•	3.33 (2.06, 5.40)	118	Unclear	Unclear	Low RoB
SQ-MSP Frozen 76-80 and 84-87 >23% 2.17 (1.38, 3.44) 108 High RoB Unclear Low RoB Yoshioka 2018 SQ-MSP Frozen 76-80 and 84-87 >2 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >2 2.18 (1.20, 3.97) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >4 36.40 1.85 (107, 3.18) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >6 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >6 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >8 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >8 1.71 (1.00, 2.93) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >8 1.71 (1.00, 2	SQ-MSP	FFPE	76-80 and 84-87	>23%				4.17 (2.35, 7.38)	118	High RoB	Unclear	Low RoB
Yoshioka 2018 SQ-MSP Frozen 76-80 and 84-87 >0 SQ-MSP Frozen 76-80 and 84-87 >2 2.72 (1.28, 5.74) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >4 1.85 (1.07, 3.18) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >6 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >6 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >6 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >6 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >8 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >8 1.71 (1.00, 2.93) 80 Low RoB Unclear Low RoB	SQ-MSP	Frozen	76-80 and 84-87	>13%				2.86 (1.79, 4.57)	108	High RoB	Unclear	Low RoB
SQ-MSP Frozen 76-80 and 84-87 >0 SQ-MSP Frozen 76-80 and 84-87 >0 SQ-MSP Frozen 76-80 and 84-87 >2 SQ-MSP Frozen 76-80 and 84-87 >4 SQ-MSP Frozen 76-80 and 84-87 >4 SQ-MSP Frozen 76-80 and 84-87 >4 SQ-MSP Frozen 76-80 and 84-87 >6 SQ-MSP Frozen 76-80 and 84-87 >6 SQ-MSP Frozen 76-80 and 84-87 >6 SQ-MSP Frozen 76-80 and 84-87 >8	SQ-MSP	Frozen	76-80 and 84-87	>23%		•	-	2.17 (1.38, 3.44)	108	High RoB	Unclear	Low RoB
SQ-MSP Frozen 76-80 and 84-87 >2 SQ-MSP Frozen 76-80 and 84-87 >4 SQ-MSP Frozen 76-80 and 84-87 >4 SQ-MSP Frozen 76-80 and 84-87 >6 SQ-MSP Frozen 76-80 and 84-87 >6 SQ-MSP Frozen 76-80 and 84-87 >8	Yoshioka 2018											
SQ-MSP Frozen 76-80 and 84-87 >2 SQ-MSP Frozen 76-80 and 84-87 >4 SQ-MSP Frozen 76-80 and 84-87 >4 SQ-MSP Frozen 76-80 and 84-87 >6 SQ-MSP Frozen 76-80 and 84-87 >6 SQ-MSP Frozen 76-80 and 84-87 >8	SQ-MSP	Frozen	76-80 and 84-87	>0	I	—		2.72 (1.28, 5.74)	80	Low RoB	Unclear	Low RoB
SQ-MSP Frozen 76-80 and 84-87 >6 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >8 1.71 (1.00, 2.93) 80 Low RoB Unclear Low RoB	SQ-MSP	Frozen	76-80 and 84-87	>2	<u> </u>	•					Unclear	Low RoB
SQ-MSP Frozen 76-80 and 84-87 >6 1.83 (1.10, 3.04) 80 Low RoB Unclear Low RoB SQ-MSP Frozen 76-80 and 84-87 >8 1.71 (1.00, 2.93) 80 Low RoB Unclear Low RoB Image: SQ-MSP Frozen 76-80 and 84-87 >8 1.71 (1.00, 2.93) 80 Low RoB Unclear Low RoB	SQ-MSP	Frozen	76-80 and 84-87	>4	+			1.85 (1.07, 3.18)	80	Low RoB	Unclear	Low RoB
SQ-MSP Frozen 76-80 and 84-87 >8 1.71 (1.00, 2.93) 80 Low RoB Unclear Low RoB	SQ-MSP	Frozen	76-80 and 84-87	>6	I ── ◆					Low RoB	Unclear	Low RoB
	SQ-MSP	Frozen		>8	+			1.71 (1.00, 2.93)	80	Low RoB	Unclear	Low RoB
\cdot						-						
				.5	1	2	4					

Comparing 5'-cytosine-phosphate-guanine-3' sites, thresholds and sample types

MSP was almost always studied in relation to CpG sites 76 to 80 and 84 to 87, so comparisons of CpG sites for MSP could not be made. One study compared MSP in FFPE versus frozen samples, and observed it to be more prognostic in frozen samples (Lattanzio 2015; see Figure 8). Many variants of PSQ were used across the studies. In particular, Dunn 2009 compared six threshold definitions for CpG sites 72 to 83; Havik 2012 compared five thresholds for CpGs 74 to 76 and four for CpGs 78 to 79; Karayan-Tapon 2010 compared six CpG sites; Quillien 2014 (test) compared 32 combinations of CpG sites and thresholds; and Quillien 2016 compared two CpG sites with multiple thresholds and both FFPE and frozen tumour samples. Figure 12 illustrates the 80 HRs for PSQ variants that we extracted or computed from 11 studies.

Figure 12. Hazard ratios from studies comparing different methods for PSQ. CpG: 5'-cytosine-phosphateguanine-3'; CI: confidence interval; FFPE: formalin-fixed paraffin embedded; HR: hazard ratio; NR: not reported; PSQ: pyrosequencing; RoB: risk of bias; sel. rep.: selective reporting.

Study ID and Sample type	CpGs analysed (PCR-based tests)	Threshold for methylated		Hazard ratio (95% CI)	Sample size	D4: PF	D5: attrition	D7: sel. rep.
Barbagallo 2014 FFPE FFPE	NR NR	>9% >25%	→	3.73 (1.68, 8.28) 1.99 (0.92, 4.32)	37 37	Law RaB Law RaB	Low RoB Low RoB	Low RoB Low RoB
Brigliadori 2016 FFPE FFPE	74-83 74-83	>9% >29%		1.92 (1.17, 3.14) 3.02 (1.72, 5.29)	94 94	Law RaB Law RaB	Law RaB Law RaB	Law RaB Law RaB
Chai 2018 (7-site cahart) Frazen Frazen Frazen	72-78 74-78 75-78	>12% >12% >12%		2.94 (1.12, 7.69) 2.94 (1.12, 7.69) 2.94 (1.12, 7.69)	24 24 24	High RaB High RaB High RaB	Law RaB Law RaB Law RaB	Unclear Unclear Unclear
Chai 2018 (8-site cohort) Frazen Frazen	75-78 75-82	> 13% > 12%		2.70 (1.37, 5.26) 3.03 (1.54, 6.25)	51 51	High RoB High RoB	Law RaB Law RaB	Unclear Unclear
Frazen Dunn 2009 Frazen, smear and/ar FFPE	76-79	>11%		2.13 (1.09, 4.17) 3.57 (2.24, 5.70)	51 109	High RoB Law RoB	Law RaB	Unclear Low RoB
Frozen, smear and/or FFPE Frozen, smear and/or FFPE Frozen, smear and/or FFPE Frozen, smear and/or FFPE Frozen, smear and/or FFPE	72-83 72-83 72-83 72-83 72-83 72-83	>20% >29% >35% Cluster 1 vs 2&3 Cluster 1&2 vs 3		4.25 (2.57, 7.05) 4.03 (2.30, 7.07) 3.64 (1.99, 6.67) 4.44 (2.58, 7.66) 3.59 (2.26, 5.69)	109 109 109 109 109	Unclear High RoB Unclear Unclear Unclear	Law RaB Law RaB Law RaB Law RaB Law RaB	Low RoB Low RoB Low RoB Low RoB Low RoB
Havik 2012 Frazen Frazen	74-78 74-78	>2.68% >6%		1.85 (1.02, 3.35) 2.30 (1.21, 4.38)	58 48	Law RaB High RaB	Low RoB	Low RoB
Frazen Frazen	74-78 74-78	>7% >8%		2.33 (1.19, 4.57) 2.33 (1.19, 4.57)	48 48	High RoB High RoB	Law RaB Law RaB	Low RoB Low RoB
Frazen Frazen	74-78 76-79	>9% >6%		2.30 (0.99, 5.33) 2.30 (1.21, 4.38)	48 48	High RoB High RoB	Law RoB Law RoB	Low RoB Low RoB
Frazen Frazen Frazen	76-79 76-79 76-79	>7% >8% >9%		2.33 (1.19, 4.57) 1.90 (0.99, 3.65) 1.90 (0.99, 3.65)	48 48 48	High RaB High RaB High RaB	Law RaB Law RaB Law RaB	Law RaB Law RaB Law RaB
Karayan-Tapon 2010 Frozen	74	>5.5%		3.26 (1.91, 5.59)	79	Low RoB	Low RoB	Low RoB
Frazen Frazen	74-78 75	>8.0% >8.7%		3.35 (1.95, 5.73) 3.26 (1.91, 5.57)	79 79	Law RoB Law RoB	Law RoB Law RoB	Low RoB Low RoB
Frazen Frazen Frazen	76 77 78	>8.0% >7.85% >7.8%		2.78 (1.64, 4.74) 3.65 (2.10, 6.34) 2.70 (1.59, 4.58)	79 79 79	Law RaB Law RaB Law RaB	Law RoB Law RoB Law RoB	Low RoB Low RoB Low RoB
Lattanzio 2015 FFPE	72-80	29%		2.09 (1.09, 3.99)	45	Low RoB	Law RaB	Low RoB
Frazen	72-80	29%	_	2.25 (1.19, 4.25)	45	Law RoB	Low RoB	Low RoB
Quillien 2014 (test) Frazen	74	>4%	_	2.44 (1.51, 3.95)	99	High RoB	Low RoB	Low RoB
Frazen Frazen	74 74-78	>8% >8%	*	2.22 (1.30, 3.79) 3.13 (1.86, 5.25)	89 99	High RoB High RoB	Law RaB Law RaB	Low RoB Low RoB
Frazen	74-78	>9%		3.13 (1.81, 5.38)	89	High RaB High RaB	Low RoB	Low RoB
Frazen Frazen	74-89 75	>11% >11%		3.13 (1.84, 5.30) 3.23 (1.87, 5.55)	89 99	High RoB	Law RaB Law RaB	Low RoB Low RoB
Frazen Frazen	75-79 76	>8% >4%		2.94 (1.71, 5.06) 2.63 (1.61, 4.31)	89 99	High RoB High RoB	Law RoB Law RoB	Low RoB Low RoB
Frazen	76	>5%		2.78 (1.62, 4.75)	89	High RoB	Low RoB	Low RoB
Frazen Frazen	76-79 76-80	>8% >9%		2.86 (1.66, 4.92) 3.03 (1.74, 5.26)	89 89	High RoB High RoB	Law RaB Law RaB	Law RaB Law RaB
Frazen Frazen	77 77	>6% >7%		3.13 (1.87, 5.23) 2.78 (1.63, 4.74)	99 89	High RoB High RoB	Law RaB Law RaB	Low RoB Low RoB
Frazen	77-81	>8%		2.70 (1.59, 4.59)	89	High RoB	Low RoB	Law RaB
Frazen Frazen	78 78	>4% >5%		2.33 (1.38, 3.92) 2.50 (1.51, 4.13)	89 99	High RoB High RoB	Law RoB Law RoB	Low RoB Low RoB
Frazen Frazen	78-82 79	>9% >7%		2.86 (1.68, 4.86) 2.78 (1.61, 4.80)	89 89	High RoB High RoB	Law RaB Law RaB	Law RaB Law RaB
Frazen	79-83	>8%		2.86 (1.68, 4.86)	89	High RoB	Low RoB	Low RoB
Frazen	80 81	>4% >8%		2.58 (1.54, 4.28) 2.44 (1.44, 4.14)	89 89	High RoB High RoB	Law RoB Law RoB	Low RoB Low RoB
Frazen Frazen	82 83	>16% >10%		2.94 (1.70, 5.08) 2.78 (1.65, 4.66)	89 89	High RaB High RaB	Law RaB Law RaB	Low RoB Low RoB
Frazen	84 84-88	>9% >17%		3.23 (1.86, 5.59) 3.23 (1.86, 5.58)	89 89	High RoB	Low RoB Low RoB	Low RoB Low RoB
Frazen	84-89	>22%		3.23 (1.85, 5.62)	89	High RoB High RoB	Low RoB	Low RoB
Frazen Frazen	85 85-89	>5% >13%		2.50 (1.52, 4.11) 2.94 (1.76, 4.92)	89 89	High RoB High RoB	Low RoB Low RoB	Low RoB Low RoB
Frazen Frazen	86 87	>11% >25%		2.78 (1.65, 4.69) 3.03 (1.75, 5.24)	89 89	High RoB	Law RaB Law RaB	Low RoB Low RoB
Frazen Frazen	87 88 89	>25% >4% >12%		3.03 (1.75, 5.24) 2.27 (1.38, 3.75) 3.23 (1.91, 5.46)	89 89	High RaB High RaB High RaB	Low RoB Low RoB	Low RoB Low RoB
Quillien 2014 (validation) FFPE	74-78	>9%		3.70 (1.71, 8.01)	50	Law RaB	Law RaB	Low RoB
FFPE	74-78 74-78	>10% >28%		2.86 (1.42, 5.74) 2.27 (0.98, 5.29)	50 50	Law RoB Law RoB	Low RoB Low RoB	Low RoB Low RoB
Quillien 2016 FFPE	74-78	>6%		3.23 (2.02, 5.16)	121	High RoB	Unclear	Low RoB
FFPE	74-78 74-78	>8% >12%		4.00 (2.30, 6.97) 4.17 (2.35, 7.38)	121 121	Low RoB Unclear	Unclear Unclear	Low RoB Low RoB
FFPE	74-78	>13%		4.35 (2.41, 7.83)	121	Unclear	Unclear	Law RaB
FFPE Frazen	74-78 74-78	>16% >6%		4.55 (2.48, 8.34) 4.00 (2.30, 6.97)	121 112	High RoB High RoB	Unclear Unclear	Low RoB Low RoB
Frazen Frazen	74-78 74-78	>8% >12 or 13%		3.57 (2.14, 5.95) 3.45 (2.10, 5.86)	112 112	Low RoB Unclear	Unclear Unclear	Low RoB Low RoB
Frazen	74-78	>12%		3.70 (2.19, 6.26)	102	Law RoB	Unclear	Low RoB
Frazen Frazen	74-78 76-79	>16% >8%		3.13 (1.94, 5.04) 3.33 (2.06, 5.40)	112 102	High RaB Law RaB	Unclear Unclear	Low RoB Low RoB
Frazen	76-79	>12%		3.33 (2.06, 5.40)	102	Low RoB	Unclear	Low RoB
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The range of CpG sites examined in studies comparing PSQ variants was from 72 to 89 (although in a comparison against IHC, Lalezari 2013 had examined sites 72 to 95). The most commonly examined CpG sites using PSQ were 74 to 78. Cut-off thresholds used when a single CpG site was targeted ranged from 4% to 25%, and

when multiple CpG sites were targeted thresholds ranged from 2.68% to 35%. In frozen tissue, Havik 2012 and Quillien 2014 (test) observed no clear dependency of HRs on threshold for CpGs 74 to 78, although Quillien 2016 observed a slight worsening as cut-off thresholds increased from 6% to 8% to 12%. In FFPE, Quillien 2014

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(validation) observed best prognostic value using a 9% threshold, followed by 10%, followed by 28% (a validation study with low risk of bias), although Brigliadori 2016 (a good-quality study) observed the opposite when comparing 9% with 29%. In a third study comparing thresholds that was free of high risks of bias, findings echoed Quillien 2014 (validation), with a threshold of 9% more prognostic than a threshold of 25%, although it was unclear what CpG islands had been targeted (Barbagallo 2014).

Quillien 2014 (test) examined a large variety of combinations of CpG sites and thresholds. Most of their highest HRs were observed for scenarios in which multiple CpG sites were targeted. There was no clear difference apparent between using PSQ on FFPE versus frozen tissue (Lattanzio 2015; Quillien 2016).

Other techniques

Bead array, PCR-mRNA and MS-MLPA

Results for studies that had examined bead array (Bady 2012 (E-GBM); Bady 2012 (M-GBM)), PCR-mRNA (Felsberg 2009; Karayan-Tapon 2010) or MS-MLPA (Park 2011) are illustrated in Figure 13. These had mostly been examined in frozen samples. One study compared two approaches to bead array with one approach to PSQ and found the highest HR for one of the bead array methods (CpG sites 76 to 84, using a cut-off threshold of 10%) with an HR of 5.56 (compared with 2.20 for PSQ). However, this particular bead array result was assessed to be at high risk of bias due to selection of the threshold. Across the evidence base, there was no clear evidence that bead array or PCR for mRNA expression differed in any consistent direction from MSP or PSQ. The only study to report MS-MLPA found it to be less prognostic than MSP (Park 2011).

Figure 13. Hazard ratios from studies comparing bead array, mRNA or MS-MLPA against either MSP or PSQ. CI: confidence interval; CpG: 5'-cytosine-phosphate-guanine-3'; FFPE: formalin-fixed paraffin embedded; HR: hazard ratio; mRNA: messenger ribonucleic acid; MS-MLPA: methylation-specific multiplex ligation-dependent probe amplification; MSP: methylation-specific polymerase chain reaction; NR: not reported; PCR: polymerase chain reaction; PF: prognostic factor; PSQ: pyrosequencing; qMSP: quantitative methylation-specific polymerase chain reaction; RoB: risk of bias; sel. rep.: selective reporting; SQ-MSP: semi quantitative methylation-specific polymerase chain reaction.

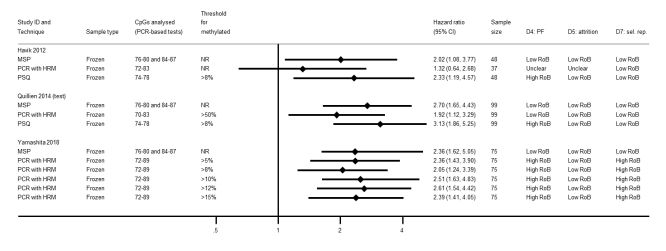
Study ID and Technique	Sample type	CpGs analysed (PCR-based tests)	Threshold for methylated		Hazard ratio (95% Cl)	Sample size	D4: PF	D5: attrition	D7: sel. rep.
Bady 2012 (E	-GBM)								
Bead array	Frozen	31 and 83	>0.358		3.28 (1.68, 6.41)	50	Low RoB	Low RoB	Low RoB
Bead array	Frozen	78-84	>10%	•	5.56 (1.25, 25.00)	50	High RoB	Low RoB	Low RoB
PSQ	Frozen	74-78	>7.28%	──	2.20 (1.12, 4.31)	47	Low RoB	Low RoB	Low RoB
Bady 2012 (M	I-GBM)								
Bead array	Frozen	31 and 83	>0.358	I ↓	6.46 (2.41, 17.35)	31	High RoB	Low RoB	Low RoB
MSP	NR	76-80 and 84-86	NR		7.21 (2.37, 21.99)	31	Low RoB	Low RoB	Low RoB
Felsberg 2009	9								
MSP	Frozen (14 FFPE)	NR	NR	──	2.23 (1.29, 3.84)	66	Low RoB	Low RoB	Low RoB
PCR-mRNA	Frozen (14 FFPE)	N/A	<50%	•	2.66 (0.94, 7.53)	23	Unclear	Unclear	Low RoB
Karayan-Tapo	on 2010								
MSP	Frozen	76-80 and 84-87	NR	──	2.32 (1.39, 3.87)	81	Low RoB	Low RoB	Low RoB
PCR-mRNA	Frozen	N/A	<0.39	→	1.68 (1.04, 2.73)	80	Low RoB	Low RoB	Low RoB
PSQ	Frozen	74-78	>8.0%	│ • ─ ─	3.35 (1.95, 5.73)	79	Low RoB	Low RoB	Low RoB
Park 2011									
MS-MLPA	50% Frozen and 50% FFPE	NR	>0.1%	── ◆───	2.38 (1.11, 5.10)	48	High RoB	Low RoB	Low RoB
MS-MLPA	50% Frozen and 50% FFPE	NR	>0.2	├── ◆───	1.88 (0.86, 4.11)	48	High RoB	Low RoB	Low RoB
MSP	50% Frozen and 50% FFPE	76-80 and 84-86	NR	│ ──◆	4.53 (1.58, 12.95)	48	Low RoB	Low RoB	Low RoB
			.5	1 2 4					

PCR-HRM

Results for three studies that had examined PCR-HRM against either MSP or PSQ (mostly in frozen samples) are presented in Figure 14. Two studies observed it to be less prognostic than MSP (Havik

2012; Quillien 2014 (test)), although a third study obtained HRs for PCR-HRM using different thresholds that fell both sides of a result for MSP, although MSP had targeted different CpG sites (Yamashita 2018).

Figure 14. Hazard ratios from studies comparing PCR-HRM against either MSP or PSQ. PCR-HRM: polymerase chain reaction with high-resolution melting; MSP: methylation-specific polymerase chain reaction; PSQ: pyrosequencing; FFPE: formalin-fixed paraffin embedded; NR: not reported; N/A: not applicable; CpG: 5'-cytosine-phosphate-guanine-3'; CI: confidence interval. RoB: risk of bias; sel. rep.: selective reporting; HRs: hazard ratios.



Further techniques

Figure 15 provides HRs for other techniques. Two linked studies (at low risk of bias) examined DIF and found it provided a better

prognostic factor than PSQ targeting CpG sites 74 to 78 in FFPE (Dahlrot 2018 (NS cohort); Dahlrot 2018 (RSD cohort)).Thon 2017 observed no difference between PSQ and sequencing.

Figure 15. Hazard ratios from of studies involving other techniques not included in previous forest plots. CI: confidence interval; CpG: 5'-cytosine-phosphate-guanine-3'; DIF: double immunofluorescence; FFPE: formalin-fixed paraffin embedded; HR: hazard ratio; IHC: immunohistochemistry; MS-RE-qPCR: methylation-specific restriction enzyme quantitative polymerase chain reaction; MSP: methylation-specific polymerase chain reaction; PSQ: pyrosequencing; QF-IHC: quantitative fluorescence immunohistochemistry; qMSP-PSQ: quantitative methylation-specific polymerase chain reaction; NR: not reported; N/A: not applicable; RoB: risk of bias; sel. rep.: selective reporting.

Study ID and Technique	Sample type	CpGs analysed (PCR-based tests)	Threshold for methylated		Hazard ratio (95% Cl)	Sample size	D4: PF	D5: attrition	D7: sel. rep
Almuqate 2018									
MS-RE-qPCR	NR	NR	>5%	• • • • • • • • • • • • • • • • • • •	4.17 (1.78, 9.75)	NR	Unclear	Unclear	Unclear
MS-RE-qPCR	NR	NR	>9%	│ • ─ • ─ • ─ • ─ • ─ • ─ • ─ • ─ •	3.57 (1.67, 7.62)	NR	Low RoB	Unclear	Unclear
Barault 2015									
Methyl-Beaming	FFPE	79-83	>40.2%	→	2.78 (1.85, 5.26)	69	Low RoB	Low RoB	Unclear
PSQ	FFPE	76-81	>29.6%	│ — ◆ — ─	2.63 (1.43, 4.55)	58	Low RoB	Unclear	Unclear
Bell 2017									
QF-IHC (AQUA)	FFPE	N/A	>Median	→	1.84 (1.38, 2.43)	452	Unclear	Low RoB	Unclear
qMSP	NR	NR	>8	→	1.77 (1.28, 2.44)	320	Unclear	Unclear	Unclear
Dahlrot 2018 (NS coh	ort)								
DIF	FFPE	N/A	<0.2	↓	1.60 (0.95, 2.71)	65	Low RoB	Low RoB	Low RoB
PSQ	FFPE	74-78	>9%	↓ • • • • • • • • • • • • • • • • • • •	1.42 (0.84, 2.40)	65	Low RoB	Low RoB	Low RoB
Dahlrot 2018 (RSD co	hort)								
DIF	FFPE	N/A	<0.2	→ _	2.00 (1.32, 3.02)	92	Low RoB	Low RoB	Low RoB
PSQ	FFPE	74-78	>10%		1.58 (1.14, 2.19)	92	Low RoB	Low RoB	Low RoB
Kristensen 2016									
IHC	FFPE	N/A	at 0%	→	1.58 (1.12, 2.22)	148	Unclear	Low RoB	Low RoB
PSQ	Frozen	NR	>10%	 →	1.80 (1.23, 2.62)	132	Low RoB	Unclear	Low RoB
qMSP-PSQ	Frozen	NR	>0.1%	→	1.64 (1.15, 2.33)	151	Low RoB	Low RoB	Low RoB
qMSP-PSQ	Frozen	NR	>5%	↓	1.66 (1.02, 2.71)	151	Low RoB	Low RoB	Low RoB
qMSP-PSQ	Frozen	NR	>20%	↓	1.52 (0.77, 3.00)	151	Low RoB	Low RoB	Low RoB
Thon 2017									
MSP	Frozen	76-80 and 84-87	NR	I ↓	3.33 (1.82, 6.25)	56	Low RoB	Low RoB	Low RoB
Sequencing	Frozen	75-99 (unclear)	>50%	│	3.33 (1.82, 6.25)	56	Low RoB	Low RoB	Low RoB
			.5	1 2 4					



Other prognostic variables

Seven studies provided both unadjusted and adjusted results for at least one method, either because these were both reported in the publication or because we could compute them from IPD. We only extracted results adjusted for age and extent of resection, or the closest to this that was available. A comparison of 15 instances of unadjusted and adjusted results for these studies is provided in Table 4. Most of these adjusted for age, but we were able to adjust additionally for resection in only one study. Two studies (six results) adjusted for Karnofsky performance status. Results were generally very similar across unadjusted and adjusted analyses, demonstrating that MGMT promoter methylation status remains prognostic after accounting for these other factors.

We did not perform formal analyses to investigate whether heterogeneity in HRs may have been due to age, extent of tumour resection, Karnofsky performance status, IDH status, recurrent versus first diagnosis, length of follow-up time of start of follow-up, due to the very limited replication of specific methods and large amounts of missing data for many of these study characteristics.

Studies examining only a single method

Table 5 provides details of the 190 articles describing studies that presented HRs from survival analysis in people in which MGMT methylation was measured by one method, and studies that used more than one method but only MGMT methylation data from one method were used in the survival analysis.

Of the 190 articles identified, 29 described studies conducted in Italy, 21 in Germany; 20 in the US; 18 in Japan; 17 in China; 11 in South Korea; nine in France; eight each in Denmark and Spain; six in the UK; three each in India, Switzerland and Taiwan; two each in Australia, Belgium, Czech Republic and Egypt; and one each in Canada, Portugal, the Netherlands and Tunisia. In addition, 23 articles described studies performed in more than one country.

Fifty-four articles reported information about follow-up, as median in months, with a minimum median of six months and a maximum of 61 months. Of these 54 articles, 29 reported data on the range of follow-up, with a minimum follow-up of 0.2 months and a maximum of 113 months. The total number of participants among the 190 articles was 27,899, with the smallest study of six and the largest study of 1395 participants. For two articles, data on number of participants were not directly reported but IPD were included.

Sixty-two articles reported data on IDH-1 and IDH-2 mutations: in 11 articles, people were 100% wild-type (i.e. no mutation), while 47 articles reported presence of IDH mutations, with mutation proportions ranging between 0.3% and 73.4% (in one study of secondary glioblastoma). Three articles did not directly report data on IDH mutations. With regard to type of tumour, 183 articles described studies only on glioblastoma, six on mixed glioblastoma and gliosarcoma, and one on gliosarcoma only.

Ninety-four articles used MSP to measure MGMT methylation, 27 used PSQ, 22 used qMSP (real-time PCR or MethyLight), 10 used bead array, four used MS-MLPA and three used PCR-HRM. In addition, 21 articles measured MGMT protein by IHC and one article by Western blotting (WB), and in four articles measured mRNA levels. Four articles reporting data from two cohorts, used a different method for each cohort.

As anticipated in our protocol (McAleenan 2019), we did not examine the results of these studies, because comparisons of HRs across studies would not provide reliable indicators of differences between the methods.

Findings: economic issues

We identified cost estimates for the three main techniques identified in the review: PSQ, MSP and IHC. These costs are expressed in 2020 Pounds Sterling (GBP). Where necessary, costs were converted using the EPPI-Centre Cost Converter (Shemilt 2019). Costs were inclusive of staff, consumables, equipment and overheads associated with preparing the sample, running the analysis and feeding back findings.

We estimated PSQ costs from personal correspondence with H Liu, Consultant Clinical Scientist, Molecular Malignancy Laboratory, Haematopathology and Oncology Diagnostic Service (HODS), Addenbrooke's Hospital, Cambridge University Hospitals NHS Foundation Trust, UK (29 October 2020). The costs were estimated to be GBP 325 per sample. Costs for IHC were estimated from a Health Technology Assessment testing for Lynch syndrome (Snowsill 2017). The costs were GBP 214.21 per sample. Finally, the costs for MSP were derived from published costing schedule from North East Thames Regional Genetics Service, the cost was GBP 198.91 (GOSH 2018).

We produced illustrative CCRs for these three tests. The comparison of PSQ to IHC would produce a CCR of 1.5. This means that to justify the increased cost, PSQ would need to have a 50% better performance. The results above indicate supremacy of PSQ over IHC (RHR 1.36, 95% CI 1.01 to 1.84), but it was unclear whether the magnitude of the superiority of the test was great enough. The CCR for PSQ and MSP was 1.6, meaning an additional 60% better performance was required. While the results presented above suggest that PSQ may have a higher RHR (1.14, 95% CI 0.87 to 1.48), the point estimate of the CCR is above the upper limit of the CI. Finally, when comparing MSP with IHC, the CCR was lower at 1.08, meaning that IHC would only require an additional 8% better performance to justify its additional cost. The results found that the lower costing intervention MSP has better ability to predict survival (RHR 1.31, 95% CI 1.01 to 1.71) and the CCR lay within the CIs for the RHR.

DISCUSSION

Summary of the main results

We examined 32 independent cohorts (3474 people) that compared different techniques, regions, cut-offs or tumour sample types for predicting overall survival in people with glioblastoma treated with temozolomide. We found evidence through meta-analysis (of RHRs) that MSP (CpG sites 76 to 80 and 84 to 87) was more prognostic than IHC (varying thresholds). Since a large majority of MSP studies had examined CpG sites 76 to 80 and 84 to 87, we were unable to compare alternative CpG sites for MSP. We also found evidence that PSQ was more prognostic than IHC, although studies of PSQ feeding into this analysis had targeted different CpG sites. The CpG sites targeted by PSQ ranged between 72 and 95, and several studies had examined sites 74 to 78. There was a suggestion that PSQ (mainly at CpG sites 74 to 78, but with varying thresholds) was slightly more prognostic than MSP (sites 76 to 80 and 84 to 87). Many variants of PSQ have been compared, although we did not see any strong

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and consistent messages from the results. Cut-off thresholds varied substantially (from 4% to 25% for single CpG sites; and from 2.68% to 35% for multiple CpG sites). Two of the three studies with low (or unclear) risk of bias that compared different thresholds found that a 9% cut-off was more prognostic than higher cut-offs (of 28% or 29%). It appears that targeting multiple CpG sites is likely to be preferable to targeting just one. We found no evidence of superiority of qMSP over MSP or PSQ, and found a suggestion that lower cut-off thresholds for interpreting SQ-MSP in frozen tissue (for sites 76 to 80 and 84 to 87) were more prognostic for survival. There was very little evidence about other techniques (such as bead array, MS-MLPA PCR-HRM) and few suggestions that any of these outperform MSP or PSQ.

Certainty of the evidence

We rated the evidence for the comparison between MSP and IHC at moderate certainty, and the evidence for comparisons of PSQ with MSP or IHC at low certainty. For all other comparisons, we rated certainty as very low. Although risk of bias and publication bias were not major concerns for us, we downgraded many of our assessments for indirectness because there was a wide variety of different CpG sites and thresholds investigated, without systematic replications of findings using the same methods across studies. The amount of evidence was small, with only 10s or at most the low 100s of participants contributing to evidence for many of the techniques.

Strengths and weaknesses of the review

We took a systematic approach to identifying, appraising and collecting information from the evidence, and assessed risk of bias and applicability concerns using a modification of QUIPS specific to the topic of the review. To ensure comparisons of methods were fair, we restricted our attention to comparisons made within studies (i.e. on the same people and tumours).

A large variety of methods have been examined, particularly use of different CpG sites and thresholds for PSQ, as well as a mixture of use of FFPE and frozen tumour samples. There was only a small amount of direct replicability across studies, meaning that firm conclusions were difficult to draw.

We limited eligibility for the review to studies that reported HRs or data sufficient for us to estimate them. In many instances, we reconstructed time-to-event data from Kaplan-Meier curves, allowing us to include 14 studies that we would not have included otherwise. However, there was still a small number of studies that had sought to compare methods but not presented data compatible with computation of HRs, which therefore did not meet our eligibility criteria. To compare HRs statistically within studies, strong correlations between different results based on the same tumour samples need to be considered. We imputed such correlations using a rather ad hoc approach and accompanied these with sensitivity analyses. We could not see an obvious way to derive appropriate correlations between these results, a problem that a statistical simulation study might help to resolve.

We listed brief details of articles describing studies examining only one method. When writing the protocol we were unsure whether these would prove to be informative about comparisons between methods. Among the studies that compared multiple methods, we observed that HRs varied markedly across studies, and we were unwilling to make naive indirect comparisons of methods across different studies. Therefore, we do not present quantitative results for these single-method studies, and did not undergo the process of establishing which articles related to the same underlying cohorts of people.

Economic issues

The results of the economic review demonstrated a paucity of economic evidence for different MGMT strategies in prediction of overall survival. Some illustrative examples showed that more costly tests might be worthwhile. However, a full economic evaluation would be required to investigate fully the costs and consequences of each of the methods for determining for MGMT promoter methylation status. The economic analyses we presented in this review were at best illustrative at this stage, since there is a lack of data to populate the decision model required to investigate the costs and consequences more robustly. A particular limitation is that cost estimates are available for only single sources and for only three of the techniques.

There are limitations with the use of overall survival as a primary outcome in terms of economic analysis. Although survival is important to consider from the perspective of patients and clinicians, other factors such as quality of life and psychological welfare associated with a patient's diagnosis and treatment are also important. This might manifest in a variety of ways, both positive and negative. For example, a false-positive test result would not have an impact on a patient's survival but would potentially cause psychological distress associated with the finding until the diagnosis was clarified, whereas a true negative might result in a sense of relief and reassurance.

For a future model-based economic evaluation, data would be needed on the costs of all the different techniques in sufficient detail to allow readers to judge applicability of the cost data, and available from more than one source to allow uncertainty to be explored. Further information would be needed to model the consequences of providing the test information: it is currently unclear how practitioners, participants and their families might use these data. Furthermore, it would be important to know whether the test results be used to stratify treatment or follow-up, and if so, what are the consequences of these changes. For both changes in management and the consequences of those changes, the impact on the use of resources, survival and health-related quality of life would be needed. As part of this, it would be important to understand fully the costs of consequences of events such as false positive and false negatives when stratifying patients, both from an overall survival and a quality-of-life perspective. This could be derived from further evidence synthesis or primary research. Alternatively, if the goal of measuring methylation status is to help the patient and their family plan for end-of-life care, then a valuation method such as willingness to pay approach, which can capture the health and non-health impacts, could be used to place a value on tests. We are not aware of any such data and hence primary research would be needed.

Applicability of findings to clinical practice and policy

We found the evidence identified to be generally applicable to clinical practice. We included only studies in which at least 90% of people had glioma, and nearly all people were treated with temozolomide. The assessments of applicability we made in our appraisal of included studies were all either of low concerns or

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unclear due to limited information. The relevance of our findings will depend to an extent on the availability of methods at local sites. For example, although many centres do PSQ, it is not available universally. Finally, we focused on overall survival only as an outcome, so are unable to draw conclusions about using these methods to predict quality of life or progression-free survival.

Agreements and disagreements with other studies or reviews

Previous systematic reviews and meta-analyses have summarised the evidence across studies using specific techniques that MGMT promoter methylation is associated with prognosis of people with glioblastoma (Hegi 2019; Zhang 2013; Zhao 2016). This is the first systematic review to our knowledge that compares methods for categorising tumours as methylated in relation to their ability to predict survival in people with glioblastoma. The findings concur with one recent consensus review on management of glioblastoma in adults from the Society for Neuro-Oncology and the European Society of Neuro-Oncology (Wen 2020). That review noted that PSQ may provide the best stratification in terms of outcomes and argued that IHC should not be used because it is unreliable. One 2020 update to evidence-based guidelines on management of newly diagnosed glioblastoma recommends the assessment of tumour MGMT promoter methylation status as a significant predictor of survival (Farrell 2020). They provided a descriptive review of the evidence about different methods but did not provide recommendations on which one(s) to use.

AUTHORS' CONCLUSIONS

Implications for practice

Among methods for categorising MGMT promoter methylation status in people with glioblastoma treated with temozolomide, pyrosequencing (PSQ) and methylation-specific polymerase chain reaction (MSP) appear to be more prognostic for overall survival than immunohistochemistry (IHC). Strong evidence is not available to draw conclusions with confidence about the best CpG sites or cut-off thresholds for quantitative methods. MSP has been studied mainly for CpG sites 76 to 80 and 84 to 87 and PSQ at CpG sites ranging from 72 to 95. A cut-off threshold of 9% for CpG sites 74 to 78 was found to perform better than higher thresholds of 28% or 29% in two of three good-quality studies making such comparisons. We are unable to draw strong conclusions about use of frozen tissue versus formalin-fixed paraffin-embedded (FFPE) in MSP and PSQ, although one study observed that MSP was more prognostic when based on frozen tissue.

Implications for research

Further large studies would be needed to establish which of the techniques has the best predictive value, and which CpG sites should be tested. Furthermore, remains important to identify a cut-off value to call the MGMT promoter "methylated," or "unmethylated," or determine ranges that predict response to temozolomide in a more graded manner. Future research should focus on MSP and PSQ (rather than IHC) as well as other up-and-coming technologies in the methylation array analysis, such as bead chip arrays.

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Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)

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* Indicates the major publication for the study

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Almuqate 2018

Study characteristics	
Study design	Cohort
Study setting	Setting: University of Calgary
	Country: Canada
	Dates: tumour samples were tested for MGMT methylation in 2015 and 2016.
Selection of participants	Cases were retrieved from the Molecular Diagnostic Laboratory database.
Participant characteristics	Sample size: 158 (deaths: NR)
	Age: mean 61 years
	Sex: 53.8% men
	KPS: NR
Tumour characteristics	GBM: 100%
	First diagnosis: NR
	Biopsy: 8.9%; subtotal resection: 38%; total resection: 53.2%
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	NR
MGMT promoter methyla- tion tests implemented	MS-RE-qPCR
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: NR; follow-up: median NR; range NR
Notes	

Bady 2012 (E-GBM)

Study characteristics	
Study design	Cohort
Study setting	Setting: Neurosurgery Departments of Rennes and Angers University Hospitals
	Country: France
	Dates: NR
Selection of participants	Cases from an external dataset (Etcheverry 2010). Prospectively collected samples from people with newly diagnosed GBM

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Bady 2012 (E-GBM) (Continued)

Participant characteristics	Sample size: 50 (deaths: NR)
	Age: median 57.5, SD NR; range 26–80 years
	Sex: 51% men
	KPS: median 78.6; range 40–100
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: NR; subtotal resection: NR; total resection: NR
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Stupp protocol
MGMT promoter methyla- tion tests implemented	PSQ, bead array
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: NR; follow-up: median NR; range NR
Notes	

Bady 2012 (M-GBM)

Study characteristics	
Study design	Cohort
Study setting	Setting: Neurosurgery Departments of Rennes and Angers University Hospitals
	Country: France
	Dates: NR
Selection of participants	Cases from an external dataset (Etcheverry 2010). Prospectively collected samples from people with newly diagnosed GBM
Participant characteristics	Sample size: 50 (deaths: NR)
	Age: median 57.5, SD NR; range 26–80 years
	Sex: 51% men
	KPS: median 78.6, range 40–100
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: NR; subtotal resection: NR; total resection: NR
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Stupp protocol

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 47

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Bady 2012 (M-GBM) (Continued)

MGMT promoter methyla- tion tests implemented	PSQ, bead array
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: NR; follow-up: median NR; range NR
Notes	

Barault 2015

Study characteristics	
Study design	Cohort
Study setting	Setting: VU University Medical Center, Amsterdam
	Country: the Netherlands
	Dates: diagnosis between 2005 and 2011
Selection of participants	Eligible people had a histopathological diagnosis of supratentorial GBM. The GBM validation-set consisted of tissue samples from people with newly diagnosed GBM, who had surgery and chemoradiation with follow-up \geq 2 years.
	Inclusion criteria: adults aged > 17 years; a new histopathological diagnosis of supratentorial GBM be- tween 2005 and 2011, verified by an independent neuropathologist; no prior brain tumour treatment to exclude dedifferentiated glioma; pre- and postoperative MRI within 3 days of surgery; standard adju- vant therapy
Participant characteristics	Sample size: 66 (deaths: NR)
	Age: NR
	Sex: % men NR
	KPS: NR
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 3%; subtotal resection: NR; total resection: 97%
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Standard adjuvant therapy consisting of radiotherapy and concomitant TMZ, followed by 6 monthly cy- cles of adjuvant TMZ
MGMT promoter methyla- tion tests implemented	PSQ, methyl-beaming
Dates and follow-up	Timing of MGMT assessment: not explicitly reported, but presumably on material obtained during re- sective surgery, prior to adjuvant therapy with 30 × 2 Gy radiotherapy and concomitant TMZ, followed by 6 monthly cycles of adjuvant TMZ.
	Start time for follow-up: NR; follow-up: median NR; range NR

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 48 people with glioblastoma treated with temozolomide (Review)

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Barault 2015 (Continued)

Notes

Study characteristics	
Study design	Cohort
Study setting	Setting: Department of Neurosurgery, Policlinico "G. Rodolico" Policlinico "G. Rodolico" University Hos pital, University of Catania
	Country: Italy
	Dates: surgery between 2004 and 2012
Selection of participants	All people underwent surgery for primary GBM with the aid of neuronavigation, and all but 2 people re- ceived gross tumour resection. The study was aimed at comparing short-term vs long-term TMZ treat- ment (people who received > 6 cycles), therefore some data were given as group A vs group B based on duration of treatment.
Participant characteristics	Sample size: 37 (deaths: NR)
	Age: mean 60.4, SD 11.8; range 30–82 years
	Sex: 51.4% men
	KPS: mean 67.1, SD 15.2
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 0%; subtotal resection: 5.4%; total resection: 94.6%
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Stupp protocol. For adjuvant TMZ therapy, people in Group A received TMZ every 28 days for > 6 cycles (up to 101), those in Group B were treated with the same adjuvant TMZ dose regimen for ≤ 6 cycles.
MGMT promoter methyla- tion tests implemented	MSP, PSQ
Dates and follow-up	Timing of MGMT assessment: NR
	Start time for follow-up: date of surgery; follow-up: median NR; range NR
Notes	

Bell 2017

Study characteristics	
Study design	Cohort nested within RCT
Study setting	Setting: NR

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 49 people with glioblastoma treated with temozolomide (Review)

Sell 2017 (Continued)	
	Country: multiple (Northern America and European centres)
	Dates: recruitment between 2006 and 2008; follow-up to 2011
Selection of participants	Participants were a subset of the NRG Oncology 0525 cohort (Gilbert 2013) with available specimens. This phase 3 trial compared standard adjuvant TMZ with a dose-dense schedule in people with newly diagnosed GBM and KPS ≥ 60.
Participant characteristics	Sample size: 452 (deaths: NR)
	Age: NR
	Sex: % men NR
	KPS: NR
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: NR; subtotal resection: NR; total resection: NR
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Standard TMZ or dose-dense TMZ
MGMT promoter methyla- tion tests implemented	qMSP, QF-IHC
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: date of randomisation, i.e. after the initial 6 weeks of chemoradiotherapy; fol- low-up: median NR; range NR
Notes	

Brigliadori 2016

Study characteristics	
Study design	Cohort
Study setting	Setting: Oncology Network of Romagna
	Country: Italy
	Dates: tumour samples collected between 2008 and 2013
Selection of participants	Participants had GBM treated with surgery and Stupp regimen. People undergoing biopsy were not in- cluded in our analysis.
Participant characteristics	Sample size: 105 (deaths: 73)
	Age: median 61, SD NR; range 23–76 years
	Sex: 61.9% men
	KPS: median NR; KPS ≤ 70: 19.1%, KPS: 80–90: 43.8%, KPS 100: 37.1%

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)

Brigliadori 2016 (Continued)

Tumour characteristics	GBM: 100%
	First diagnosis: NR
	Biopsy: 0%; subtotal resection: 51.1%; total resection: 49%
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Stupp protocol
MGMT promoter methyla- tion tests implemented	PSQ, Bead array
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: date of diagnosis; follow-up: median 55; range 5–79 months
Notes	

Chai 2018 (7-site cohort)

Study characteristics	
Study design	Cohort
Study setting	Setting: Chinese Glioma Genome Atlas (CGGA) database
	Country: China
	Dates: NR
Selection of participants	Inclusion criteria: diagnosed with WHO grade III or IV glioma; containing MGMT promoter methylation PSQ testing data in detail; including exact MGMT mRNA sequencing data; having received radiotherapy + TMZ treatment; containing overall survival information
Participant characteristics	Sample size: 24 (deaths: NR)
	Age: median 55; range 29–79 years
	Sex: 58.3% men
	KPS: median NR; KPS < 80: 11/24 (45.8%); KPS ≥ 80: 9/24 (37.5%); KPS not available: 4/24 (16.7%)
Tumour characteristics	GBM: 100%
	First diagnosis: NR
	Biopsy: 0%; subtotal resection: 50%; total resection: 50%
	IDH mutant 5/24 (20.8%), IDH wild-type 18/24 (75.0%), not available 1/24 (4.2%). IDH status combined results of IDH1 and IDH2 testing
Treatment regimen	Radiotherapy + TMZ
MGMT promoter methyla- tion tests implemented	PSQ
Dates and follow-up	Timing of MGMT assessment: not explicitly reported, but presumably on freshly frozen tumour samples obtained during resection/biopsy.

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)



Chai 2018 (7-site cohort) (Continued)

Start time for follow-up: NR; follow-up: median NR; range NR

Notes

Chai 2018 (8-site cohort)

Study characteristics	
Study design	Cohort
Study setting	Setting: Chinese Glioma Genome Atlas (CGGA) database
	Country: China
	Dates: NR
Selection of participants	Inclusion criteria: diagnosed with WHO grade III or IV glioma; containing MGMT promoter methylation PSQ testing data in detail; including exact MGMT mRNA sequencing data; having received radiotherapy + TMZ treatment; containing overall survival information
Participant characteristics	Sample size: 24 (deaths: NR)
	Age: median 55; range 29–79 years
	Sex: 58.3% men
	KPS: median NR; KPS < 80: 11/24 (45.8%); KPS ≥ 80: 9/24 (37.5%); KPS not available: 4/24 (16.7%)
Tumour characteristics	GBM: 100%
	First diagnosis: NR
	Biopsy: 0%; subtotal resection: 50%; total resection: 50%
	IDH mutant 5/24 (20.8%), IDH wild-type 18/24 (75.0%), not available 1/24 (4.2%). IDH status combined results of IDH1 and IDH2 testing.
Treatment regimen	Radiotherapy + TMZ
MGMT promoter methyla- tion tests implemented	PSQ
Dates and follow-up	Timing of MGMT assessment: not explicitly reported, but presumably on freshly frozen tumour samples obtained during resection/biopsy.

Dahlrot 2018 (NS cohort)

Study characteristics	
Study design	Cohort
Study setting	Setting: Nordic study (validation cohort)

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 52

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Dahlrot 2018 (NS cohort) (Continued) Country: Denmark

	, ,
	Dates: diagnosis between January 2003 and May 2008.
Selection of participants	People included in a collaborative Nordic Study (NS) with WHO grade 3 and 4 gliomas treated with ra- diotherapy and different combinations of TMZ. Minimum 15 mm ² vital tumour tissue was required for inclusion.
Participant characteristics	Sample size: 92 (deaths: 64)
	Age: NR
	Sex: 57% men
	KPS: median NR ECOG performance status: 0–1: 64 (94%); 2: 4 (6%)
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: NR; subtotal resection: NR; total resection: NR
	IDH1 wild-type: 94%; IDH2 wild-type: NR
Treatment regimen	Radiotherapy and different combinations of TMZ
MGMT promoter methyla- tion tests implemented	PSQ, DIF
Dates and follow-up	Timing of MGMT assessment: NR
	Start time for follow-up: date of randomisation; follow-up: median 17.5; range 0.5–129 months
Notes	

Dahlrot 2018 (RSD cohort)

Study characteristics	
Study design	Cohort
Study setting	Setting: Region of Southern Denmark
	Country: Denmark
	Dates: diagnosis between 1 January 2005 to 31 December 2009
Selection of participants	Inhabitants in the Region of Southern Denmark (RSD), and no treatment received prior to surgery. Mini- mum 15 mm ² vital tumour tissue required for inclusion.
Participant characteristics	Sample size: 234 (deaths: 168)
	Age: median NR; aged < 65 years: 83 (49%); aged > 65 years: 88 (51%)
	Sex: 57% men
	KPS: median NR; ECOG performance status: 0–1: 106 (62%); 2–4: 65 (38%)
Tumour characteristics	GBM: 100%

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 53 people with glioblastoma treated with temozolomide (Review)

Dahlrot 2018 (RSD cohort) (Continued)

	First diagnosis: NR
	Biopsy: NR; subtotal resection: NR; total resection: NR
	IDH1 wild-type: 98%; IDH2 wild-type: NR
Treatment regimen	Stupp regimen (57%); palliative (25%); none (18%)
MGMT promoter methyla- tion tests implemented	PSQ, DIF
Dates and follow-up	Timing of MGMT assessment: NR
	Start time for follow-up: date of primary surgery; follow-up: median 11; range 0.03–96 months
Notes	

Dunn 2009

Study characteristics	
Study design	Cohort
Study setting	Setting: surgery at Walton Centre for Neurology and Neurosurgery and treatment at Clatterbridge Cen- tre for Oncology
	Country: UK
	Dates: diagnosis between June 2004 and October 2007
Selection of participants	Newly diagnosed, previously untreated GBMs WHO grade IV. These people had cytoreductive surgery where possible or biopsy before radical treatment with radiotherapy and concurrent TMZ + radiotherapy py followed 4 weeks later by adjuvant TMZ.
Participant characteristics	Sample size: 109 (deaths: 94)
	Age: median 55; range 18–68 years
	Sex: 66.1% men
	KPS: median NR; WHO performance status 0: 37/109 (33.9%); WHO performance status 1: 54/109 (49.5%); WHO performance status 2: 16/109 (14.7%); WHO performance status 3: 2/109 (1.8%)
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 23.9%; subtotal resection: 0%; total resection: 76.1%; dichotomised as biopsy or resection
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Radiotherapy and concurrent TMZ + radiotherapy followed 4 weeks later by adjuvant TMZ
MGMT promoter methyla- tion tests implemented	PSQ
Dates and follow-up	Timing of MGMT assessment: not explicitly reported, but presumably on freshly frozen tumour samples obtained during resection/biopsy

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 54 people with glioblastoma treated with temozolomide (Review)



Dunn 2009 (Continued)

Start time for follow-up: date of diagnosis; follow-up: median NR; range NR

Notes

Study characteristics	
Study design	Cohort
Study setting	Setting: Department of Neurosurgery, Heinrich-Heine-University Düsseldorf
	Country: Germany
	Dates: recruitment between 1998 and 2004; follow-up to 2006
Selection of participants	Participants had been treated with open resection and ≥ 2 cycles of chemotherapy with TMZ first-line. They had sufficient tissue for molecular analysis, and available follow-up data.
Participant characteristics	Sample size: 67 (deaths: 58)
	Age: median 56; range 26–80 years
	Sex: 61.2% men
	KPS: median 80; range 20–90
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 0%; subtotal resection: NR; total resection: NR; NCR defined by a residual tumour volume of < 5 mL on early postoperative MRI
	IDH1 wild-type: 100%; IDH2 wild-type: NR
Treatment regimen	Radiotherapy followed by adjuvant TMZ according to the standard 5-day schedule every 28 days
MGMT promoter methyla- tion tests implemented	MSP, IHC, PCR-mRNA
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: date of surgery for primary tumour; follow-up: median 40.6; range 16.5–96.0 months

Havik 2012

Study characteristics	
Study design	Cohort
Study setting	Setting: Department of Neurosurgery, Oslo University Hospital

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 55 people with glioblastoma treated with temozolomide (Review)

Havik 2012 (Continued)	
	Country: Norway
	Dates: surgery between January 2005 and January 2009
Selection of participants	Tumour samples from 134 people with glioma (diffuse astrocytoma WHO grade II (n = 10), oligoden- droglioma WHO grade II (n = 6), oligoastrocytoma WHO grade II (n = 17), low-grade neuroepithelial tu- mour not otherwise specified (n = 2), anaplastic astrocytoma WHO grade III (n = 4), anaplastic oligoden- droglioma, WHO grade III (n = 6), anaplastic oligoastrocytoma WHO grade III (n = 3), GBM WHO grade IV (n = 86)) and 4 people with meningioma
Participant characteristics	Sample size: 134 (deaths: NR)
	Age: mean 58.5, SD 9.1 years
	Sex: 53.5% men
	KPS: NR
Tumour characteristics	GBM: 64.2%
	First diagnosis: NR
	Biopsy: NR; subtotal resection: NR; total resection: NR
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Standard radiotherapy and concomitant TMZ, some also adjuvant TMZ
MGMT promoter methyla- tion tests implemented	MSP, PSQ, qMSP, PCR-HRM
Dates and follow-up	Timing of MGMT assessment: not explicitly reported, but presumably on freshly frozen tumour samples obtained during resection/biopsy.
	Start time for follow-up: date of first surgery; follow-up: median NR; range NR
Notes	

Hsu 2015

Study characteristics	
Study design	Cohort
Study setting	Setting: Taipei Veterans General Hospital
	Country: Taiwan, Republic of China
	Dates: enrolment between October 2007 and January 2014
Selection of participants	People with primary GBM, TMZ chemotherapy with concomitant radiotherapy and adequate follow-up data
Participant characteristics	Sample size: 121 (deaths: 119)
	Age: median 55; range 40–65 years
	Sex: 59.5% men

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Hsu 2015 (Continued)

	KPS: median NR; KPS ≥ 80: 66 (54.6%)
Tumour characteristics	GBM: 100%
	First diagnosis: NR
	Biopsy: NR; subtotal resection: NR; total resection: 81%
	IDH1 wild-type: 91.7%; IDH2 wild-type: NR
Treatment regimen	TMZ chemotherapy with concomitant radiotherapy
MGMT promoter methyla- tion tests implemented	MSP, PSQ, IHC, qMSP
Dates and follow-up	Timing of MGMT assessment: NR, probably after surgery
	Start time for follow-up: date of surgery; follow-up: median NR; range NR
Notes	

Karayan-Tapon 2010

Study characteristics	
Study design	Cohort
Study setting	Setting: Centres in Poitiers, Rennes and Nantes
	Country: France
	Dates: NR
Selection of participants	Participants had GBM treated with surgery and Stupp regimen
Participant characteristics	Sample size: 81 (deaths: NR)
	Age: median 61; range 30–78 years
	Sex: 55.6% men
	KPS: median NR; WHO performance status: 0–2: 52/81 (64.2%); WHO performance status 3–4: 23/81 (28.4%); WHO performance status not available: 6/81 (7.4%)
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: NR; subtotal resection: NR; total resection: NR; extent of resection determined perioperatively by neurosurgeon
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Stupp protocol
MGMT promoter methyla- tion tests implemented	MSP, PSQ, IHC, SQ-MSP, PCR-mRNA
Dates and follow-up	Timing of MGMT assessment: at diagnosis

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 57



Karayan-Tapon 2010 (Continued)

Start time for follow-up: date of surgery; follow-up: median 16; range 5–57 months

Notes

Study characteristics	
Study design	Cohort
Study setting	Setting: University College of Medicine, Busan; Ungkyunkwan University School of Medicine, Changwor
	Country: South Korea
	Dates: tissue collected between 1997 and 2012
Selection of participants	Study set included FFPE brain tumour tissue diagnosed as GBM. All people underwent surgical resec- tion or biopsy sampling of their tumours
Participant characteristics	Sample size: 104 (deaths: 79)
	Age: mean 51.4; range 26.4–87.2 years
	Sex: 55.8% men
	KPS: median NR; KPS ≥ 70: 71.2%; KPS < 70: 28.8%
Tumour characteristics	GBM: 100%
	First diagnosis: NR
	Biopsy: 4.8%; subtotal resection: 57.7%; total resection: 37.5%
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Concurrent TMZ chemoradiotherapy
MGMT promoter methyla- tion tests implemented	MSP, PSQ
Dates and follow-up	Timing of MGMT assessment: not explicitly reported but presumably on tissue harvested during biop- sy/resection
	Start time for follow-up: date of diagnosis; follow-up: median NR; range 3.2–41.5 months

Kristensen 2016

Study characteristics	
Study design	Cohort
Study setting	Setting: Rigshospitalet
	Country: Denmark

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 58 people with glioblastoma treated with temozolomide (Review)



Kristensen 2016 (Continued)	Dates: cases diagnosed between 2005 and 2010. Study ended 2015
Selection of participants	Participants had available samples and received Stupp regimen
Participant characteristics	Sample size: 151 (deaths: 146)
	Age: median 59, SD NR; range 22–74 years
	Sex: 62.9% men
	KPS: NR
Tumour characteristics	GBM: 100%
	First diagnosis: NR
	Biopsy: 3.3%; subtotal resection: NR; total resection: 41.1%; NR
	IDH1 wild-type: 96%; IDH2 wild-type: NR
Treatment regimen	Stupp protocol
MGMT promoter methyla- tion tests implemented	IHC, qMSP-PSQ
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: NR; follow-up: median 94; range 53–123 months
Notes	

Lalezari 2013

Study characteristics	
Study design	Cohort
Study setting	Setting: University of California Los Angeles and Kaiser Permanente Los Angeles
	Country: USA
	Dates: diagnosed between 2000 and 2010.
Selection of participants	People were retrospectively identified based on an electronic database query of adults with prima- ry GBM receiving upfront TMZ and treated at the University of California Los Angeles or Kaiser Perma- nente Los Angeles. People whose samples were directed to the laboratory in an unselected manner were also included.
Participant characteristics	Sample size: 418 (deaths: 356)
	Age: median 57.6; range 22.3–90.0 years
	Sex: 60.8% men
	KPS: median NR; KPS 100: 13.9%; KPS 90: 47.6%; KPS 80: 24.6%; KPS 70: 6.2%; KPS ≤ 60: 7.2%; KPS missing: 0.5%
Tumour characteristics	GBM: 100%

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 59 people with glioblastoma treated with temozolomide (Review)

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Lalezari 2013 (Continued)	First diagnosis: 100% Biopsy: 9.1%; subtotal resection: 47.8%; total resection: 41.9%; NR IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Combination of radiotherapy and TMZ: concurrent daily radiotherapy/TMZ followed by TMZ (Stupp, n = 235), maintenance dose TMZ overlapping with radiotherapy (modified Stupp, n = 127), TMZ after radio-therapy (pre-Stupp, n = 48).
MGMT promoter methyla- tion tests implemented	MSP, PSQ, IHC
Dates and follow-up	Timing of MGMT assessment: FFPE samples from initial surgery prior to any treatment
	Start time for follow-up: NR; follow-up: median 70; range 2–137 years
Notes	

Lattanzio 2015

Study characteristics

Cohort
Setting: Department of Neurosurgery, Santa Croce University Hospital, Cuneo
Country: Italy
Dates: tissue collected between 2006 and 2013
People with newly diagnosed GBM and treated with standard TMZ-containing chemoradiotherapy pro- tocols
Sample size: 46 (deaths: 29)
Age: median 64.5; range 24–84 years
Sex: 76.1% men
KPS: NR
GBM: 100%
First diagnosis: 100%
Biopsy: NR; subtotal resection: NR; total resection: NR
IDH1 wild-type: NR; IDH2 wild-type: NR
Standard TMZ-containing chemoradiotherapy protocols
MSP, PSQ
Timing of MGMT assessment: for each participant, 2 samples of the primary tumour obtained: 1 collect- ed during surgery, immersed in RNA later (Life Technologies, Carlsbad, CA, USA) and immediately snap- frozen in liquid nitrogen, and 1 assembled from biopsy in FFPE sections using standard procedures.

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)



Lattanzio 2015 (Continued)

Start time for follow-up: date of first surgery; follow-up: median 7.4 months; range NR

Notes

Lechapt-Zalcman 2012

Study characteristics	
Study design	Cohort
Study setting	Setting: network of 11 neurosurgical university departments
	Country: France
	Dates: tumour samples collected between 2005 and 2009
Selection of participants	Tumour samples collected from 2 observational studies analysing the use of Gliadel implants in people with newly diagnosed GBM
Participant characteristics	Sample size: 111 (deaths: 56)
	Age: median 58, SD NR; range 33–77 years
	Sex: 65.8% men
	KPS: mean 80.2, SD 13.5
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 0%; subtotal resection: 30%; total resection: 55.5%; total 100% disappearance of contrast enhancement, subtotal ≥ 90% disappearance of contrast enhancement, partial < 90% disappearance of contrast enhancement, partial < 90% disappearance of contrast enhancement.
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Gliadel (carmustine) wafers followed by the Stupp protocol
MGMT promoter methyla- tion tests implemented	MSP, PSQ
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: NR; follow-up: median 13.6; range 0–37.6 months
Notes	

McDonald 2013

Study characteristics	
Study design	Cohort
Study setting	Setting: Royal North Shore Hospital and the North Shore Private Hospital, Sydney

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 61 people with glioblastoma treated with temozolomide (Review)



McDonald 2013 (Continued)

	Country: Australia
	Dates: NR
Selection of participants	Retrospective cohort of people with primary GBM treated by gross total resection
Participant characteristics	Sample size: 78 (deaths: 74)
	Age: mean 58.4, SD 12.4; range 22–83 years
	Sex: 75.6% men
	KPS: NR
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 0%; subtotal resection: 0%; total resection: 100%
	IDH1 wild-type: 97.4%; IDH2 wild-type: NR
Treatment regimen	Concurrent radiotherapy and TMZ followed by adjuvant TMZ (61.5%) or TMZ as an adjuvant therapy af- ter radiotherapy (38.5%)
MGMT promoter methyla- tion tests implemented	MSP, PSQ
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: NR; follow-up: median NR; range NR
Notes	

Melguizo 2012

Study characteristics	
Study design	Cohort
Study setting	Setting: Hospital Virgen de las Nieves from Granada (Spain) and the University Hospital of Sassari (Italy)
	Country: Spain and Italy
	Dates: tumour samples collected between 2001 and 2009
Selection of participants	Participants aged \geq 70 years with newly diagnosed GBM and postoperative KPS \geq 60
Participant characteristics	Sample size: 78 (deaths: NR)
	Age: mean 56; range 24–81 years
	Sex: 53.8% men
	KPS: median NR. All participants had KPS ≥ 60
Tumour characteristics	GBM: 100%
	First diagnosis: 100%

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 62 people with glioblastoma treated with temozolomide (Review)

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	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Concurrent chemoradiotherapy with TMZ followed by adjuvant TMZ
MGMT promoter methyla- tion tests implemented	MSP, IHC
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: date of diagnosis; follow-up: median NR; range NR
Notes	

Nguyen 2015

Study characteristics	
Study design	Cohort
Study setting	Setting: Oncological Centre, University Hospital of Strasbourg
	Country: France
	Dates: cases treated and followed up between 2006 and 2010
Selection of participants	Participants aged > 18 years with treatment-naive GBM
Participant characteristics	Sample size: 106 (deaths: NR)
	Age: median NR, SD NR; aged ≥ 50 years: 78%, aged < 50 years: 22%
	Sex: 63% men
	KPS: NR
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 12%; subtotal resection: 34%; total resection: 34%; gross-total (no residual tumour on MRI), subtotal (residual tumour on MRI)
	IDH1 wild-type: 98%; IDH2 wild-type: NR
Treatment regimen	Stupp protocol
MGMT promoter methyla- tion tests implemented	FSQ-MS-PCR
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: NR; follow-up: median 17.4; range 2–92.8 months
Notes	

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in
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Park 2011

Study characteristics	
Study design	Cohort
Study setting	Setting: Seoul National University Hospital and Seoul National University Bundang Hospital
	Country: South Korea
	Dates: NR
Selection of participants	Participants with newly diagnosed supratentorial GBM treated with surgery and Stupp regimen
Participant characteristics	Sample size: 48 (deaths: 22)
	Age: mean 53.4; range 28–74 years
	Sex: 62.5% men
	KPS: NR
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 24%; subtotal resection: NR; total resection: NR
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Stupp protocol
MGMT promoter methyla- tion tests implemented	MSP, MS-MLPA
Dates and follow-up	Timing of MGMT assessment: at diagnosis
	Start time for follow-up: date of surgery; follow-up: median 16 months; range NR
Notes	

Quillien 2014 (test)

Study characteristics	
Study design	Cohort
Study setting	Setting: 4 centres (Marseille, Paris, Poitiers and Rennes)
	Country: France
	Dates: treatment between November 2003 and September 2007
Selection of participants	People with newly diagnosed primary GBM, excluding giant-cell GBM, were given standard care treat- ment and followed up for ≥ 18 months. For each participant, a frozen tumour sample and paraffin-em- bedded tissue specimens had to be available.
	People treated between November 2003 and September 2007

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 64 people with glioblastoma treated with temozolomide (Review)

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Quillien 2014 (test) (Continued)	
Participant characteristics	Sample size: 100 (deaths: 75)
	Age: median 57.5; range 21.0–73.0 years
	Sex: 64% men
	KPS: median NR; KPS 90–100: 28%; KPS 70–80: 56%; KPS < 70: 16%
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 7%; subtotal resection: 22%; total resection: 71%
	IDH1 wild-type: 100; IDH2 wild-type: NR
Treatment regimen	TMZ chemotherapy with concomitant radiotherapy, followed by cycles of adjuvant TMZ. 57 people re- quired second-line treatment (nitrosourea chemotherapy (n = 30); surgery with carmustine wafers (n = 12); surgery + nitrosourea chemotherapy (n = 1); bevacizumab + irinotecan (n = 6); other chemother- apy (n = 8). 12 people required third-line treatment (bevacizumab + irinotecan (n = 5); nitrosourea chemotherapy (n = 3); other chemotherapy (n = 4)
MGMT promoter methyla- tion tests implemented	MSP, PSQ, IHC, MethyLight-MSP, PCR-HRM
Dates and follow-up	Timing of MGMT assessment: tumour samples obtained during surgery
	Start time for follow-up: NR; follow-up: median 17.9 months; range NR
Notes	

Quillien 2014 (validation)

Study characteristics	
Study design	Cohort
Study setting	Setting: NR
	Country: France
	Dates: NR
Selection of participants	Independent validation cohort comprised 50 people with newly diagnosed GBM treated with radiother- apy and concurrent/adjuvant TMZ
Participant characteristics	Sample size: 50 (deaths: NR)
	Age: median 59; range 41–78 years
	Sex: % men NR
	KPS: median NR; KPS 90–100: 22 (44%); KPS 70–80: 23 (46%); KPS < 70: 5 (10%)
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: NR; subtotal resection: NR; total resection: NR

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 65 people with glioblastoma treated with temozolomide (Review)



Quillien 2014 (validation) (Continued)

	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Radiotherapy and concurrent/adjuvant TMZ
MGMT promoter methyla- tion tests implemented	PSQ
Dates and follow-up	Timing of MGMT assessment: NR
	Start time for follow-up: NR; follow-up: median NR; range NR
Notes	

Quillien 2016

Study characteristics	
Study design	Cohort
Study setting	Setting: 8 centres
	Country: France
	Dates: enrolled between March 2009 and June 2011
Selection of participants	Inclusion criteria: histologically confirmed de novo-GBM, aged 18–70 years, presented with no con- traindications as dictated by the Stupp protocol and not included in experimental therapeutic proto- cols
Participant characteristics	Sample size: 139 (deaths: 119)
	Age: median 55.9; range 23.0–71.0 years
	Sex: 70.5% men
	KPS: median NR; KPS 90–100: 41 (29.5%); KPS 70–80: 76 (54.7%); KPS < 70: 20 (14.4%); KPS missing: 2 (1.4%)
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 14.4%; subtotal resection: 29.5%; total resection: 56.1%
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Stupp protocol
MGMT promoter methyla- tion tests implemented	PSQ, SQ-MSP
Dates and follow-up	Timing of MGMT assessment: not explicitly reported but presumably on tissue harvested during biop- sy/resection
	Start time for follow-up: NR; follow-up: median NR; range NR

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 66 people with glioblastoma treated with temozolomide (Review)

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Thon 2017

Study characteristics	
Study design	Cohort
Study setting	Setting: University of Munich
	Country: Germany
	Dates: enrolment between March 2006 and August 2008; last follow-up June 2016
Selection of participants	Adults with supratentorial GBM not suitable for gross total tumour resection with histology being con- firmed by stereotactic biopsy; no severe mass effect of the tumour demanding debulking surgery; no prior history of surgery, radiotherapy or chemotherapy (or both); KPS ≥ 60 and adequate haematologi- cal, renal and hepatic function (Thon 2011)
Participant characteristics	Sample size: 56 (deaths: 53)
	Age: median 62.5; range 23–85 years
	Sex: 58.9% men
	KPS: median 70; inclusion criterion was KPS ≥ 60. 24 (42.9%) participants had KPS 70 and 13 had KPS 60 (23.2%)
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 100%; subtotal resection: 0%; total resection: 0%
	IDH1 wild-type: NR; IDH2 wild-type: NR
Treatment regimen	Radiotherapy/TMZ followed by adjuvant TMZ (adjuvant TMZ was not initiated in 14 people because of clinical deterioration with disorientation and confusion). Salvage treatment for progressive disease was initiated in 22 people and best supportive care in 33 people.
MGMT promoter methyla- tion tests implemented	MSP, PSQ
Dates and follow-up	Timing of MGMT assessment: tissue samples collected during biopsy
	Start time for follow-up: date of biopsy; follow-up: median NR; range NR
Notes	

Yamashita 2018

Study characteristics	
Study design	Cohort
Study setting	Setting: Miyazaki University Hospital
	Country: Japan
	Dates: surgery between February 2008 and July 2015

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Yamashita 2018 (Continued)	
Selection of participants	People with newly diagnosed GBM who had undergone surgery
Participant characteristics	Sample size: 75 (deaths: NR)
	Age: median 64; range 32–84 years
	Sex: 61.3% men
	KPS: median NR; KPS 90–100: 18 (24%); KPS 70–80: 34 (45.3%); KPS < 70: 23 (30.7%)
Tumour characteristics	GBM: 100%
	First diagnosis: 100%
	Biopsy: 2.7%; subtotal resection: 30.7%; total resection: 66.7%
	92% (IDH1 mutated 6.7%; unknown 1.3%); IDH2 wild-type: NR
Treatment regimen	Stupp protocol
MGMT promoter methyla- tion tests implemented	MSP, PCR-HRM
Dates and follow-up	Timing of MGMT assessment: tissue specimens obtained at surgery
	Start time for follow-up: NR; follow-up: median 17 months; range NR
Notes	

Yang 2012

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Study characteristics	
Study design	Cohort
Study setting	Setting: Catholic University of Korea, Suwon
	Country: South Korea
	Dates: surgery between 2000 and 2006
Selection of participants	People undergoing surgery with new histological diagnosis of supratentorial GBMs classified according to the WHO 2007 criteria
Participant characteristics	Sample size: 18 (deaths: 13)
	Age: mean 53.3, SD 14.1; range 23–71 years
	Sex: 50% men
	KPS: NR
Tumour characteristics	GBM: 100%
	First diagnosis: NR
	Biopsy: 5.6%; subtotal resection: 27.8%; total resection: 66.7%
	IDH1 wild-type: NR; IDH2 wild-type: NR

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Yang 2012 (Continued)	
Treatment regimen	Radiotherapy + TMZ (66.7%); radiotherapy + PCV + TMZ (33.3%)
MGMT promoter methyla- tion tests implemented	MSP, IHC
Dates and follow-up	Timing of MGMT assessment: tumour samples obtained during surgery Start time for follow-up: date of histological diagnosis; follow-up: median NR; range NR
Notes	

Yoshioka 2018

Study characteristics	
Study design	Cohort
Study setting	Setting: Chiba University Hospital
	Country: Japan
	Dates: NR
Selection of participants	People under a protocol approved by the Ethics Committee of the Chiba University Graduate School of Medicine, with informed consent obtained from the people or their guardians
Participant characteristics	Sample size: 84 (deaths: NR)
	Age: median NR; aged < 60 years: 36 (43%); aged ≥ 60 years: 48 (57%)
	Sex: 51% men
	KPS: median NR; KPS ≤ 70: 47 (56%); KPS > 70: 37 (44%)
Tumour characteristics	GBM: 100%
	First diagnosis: NR
	Biopsy: NR; subtotal resection: 0.57%; total resection: 0.43%
	IDH1 wild-type: 94%; IDH2 wild-type: NR
Treatment regimen	Stupp protocol
MGMT promoter methyla- tion tests implemented	MSP
Dates and follow-up	Timing of MGMT assessment: time of the first surgery
	Start time for follow-up: date of initial surgery; follow-up: median NR; range NR
Notes	

DIF: double immunofluorescence; ECOG: Eastern Cooperative Oncology Group; FFPE: formalin-fixed paraffin-embedded; FSQ-MS-PCR: fluorescent semi-quantitative methylation-specific polymerase chain reaction; GBM: glioblastoma; IDH: isocitrate dehydrogenase; IHC: immunohistochemistry; KPS: Karnofsky performance status; MGMT: O⁶-methylguanine–DNA methyltransferase; mRNA: messenger ribonucleic acid; MRI: magnetic resonance imaging; MS-MLPA: methylation-specific multiplex ligation-dependent probe amplification; MS-

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 69



RE-qPCR: methylation-specific restriction enzyme quantitative polymerase chain reaction; MSP: methylation-specific polymerase chain reaction; n: number of participants; NR: not reported; PCR: polymerase chain reaction; PCR-HRM: polymerase chain reaction with high-resolution melting; PCR-mRNA: polymerase chain reaction-messenger ribonucleic acid; PCV: procarbazine plus lomustine plus vincristine; PSQ: pyrosequencing; QF-IHC: quantitative fluorescence immunohistochemistry; qMSP: quantitative methylation-specific polymerase chain reaction; with pyrosequencing; RCT: randomised controlled trial; RNA: ribonucleic acid; SD: standard deviation; SQ-MSP: semi-quantitative methylation-specific polymerase chain reaction; TMZ: temozolomide; WHO: World Health Organization.

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Becker 2016	HR not reported/could not be calculated.
Becker 2018	Unclear if all people received TMZ.
Christians 2012	HR not reported/could not be calculated.
Crosby 2013	HR not reported/could not be calculated.
Gurrieri 2018	HR not reported/could not be calculated.
Jung 2010	People had to survive until recurrence to be included.
Kang 2011	IPD were available for the 9 eligible people, but these were too few to estimate HRs with any relia- bility.

HR: hazard ratio; IPD: individual participant data; TMZ: temozolomide.

Characteristics of studies awaiting classification [ordered by study ID]

Cao 2009

Notes	Unclear report, further information required to make a decision.	
Notes	Unclear report, further information required to make a decision.	

Dreval 2009

Notes	Requires translation (Russian).

Ellingson 2012

Notes

Unclear report, further information required to make a decision.

Fosmark 2017

Notes

Unclear report, further information required to make a decision.

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Grabenbauer 2010

Notes

Unclear report, further information required to make a decision.

Hou 2011	
Notes	Requires translation (Chinese).
Jarboe 2012	
Notes	Unclear report, further information required to make a decision.
Kalkan 2015	
Notes	Unclear report, further information required to make a decision.
Kamoshima 2012	
Notes	Requires translation (Japanese).
Lin 2008	
Notes	Requires translation (Chinese).
Liu 2018	
Notes	Requires translation (Chinese).
Lobanova 2016	
Notes	Requires translation (Russian).
Shen 2011	
Notes	Requires translation (Chinese).
Sun 2004	
Notes	Requires translation (Chinese).

people with glioblastoma treated with temozolomide (Review)



Tang 2012

Notes

Unclear report, further information required to make a decision.

Yan 2015	
Notes	Requires translation (Chinese).

Yang 2011

Notes Requires translation (Chinese).

Characteristics of ongoing studies [ordered by study ID]

Rapp 2018

Study name	GlioVax
Starting date	March 2018
Contact information	Michael Sabel (Michael.Sabel@med.uni-duesseldorf.de)
Notes	Trial identifier: EudraCT-Number 2017–000304-14
	Country: Germany
	Inclusion criteria: monofocal GBM, IDH wild-type; near-complete resection (≤ 5 mL residual tumour volume); Karnofsky performance status ≥ 70%

GBM: glioblastoma; IDH: isocitrate dehydrogenase.

ADDITIONAL TABLES

Table 1. Techniques of determining methylation status

Test	Brief description
Methylation-specific poly- merase chain reaction (MSP)	In MSP, DNA is extracted from tumour tissue and then treated with sodium bisulfite. Sodium bisul- fite causes changes in the sequence of unmethylated DNA, as it changes the DNA base cytosine into uracil. Methylated DNA is protected and remains unchanged. Regions of DNA can then be amplified using PCR in a manner that is dependent on whether the changed (containing uracil) or original se- quence (containing cytosine) is present.
Quantitative (or real time) methylation-specific PCR (qMSP)	qMSP is very similar to MSP, but there is a measure of the amount of changed and original DNA se- quence.
Methylation-specific sequenc- ing, including pyrosequencing (PSQ)	In methylation-specific sequencing, DNA is extracted from tumour tissue and treated with sodium bisulfite, which changes unmethylated DNA. The DNA can then be sequenced to determine if it con-

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 72

Table 1. Techniques of determining methylation status (Continued)

	tains the changed or original sequence, i.e. whether it contains uracil in place of cytosine. There are many ways of sequencing DNA, but one commonly used method is called PSQ.
Bead array	In bean array, DNA is extracted from tumour tissue and treated with sodium bisulfite, which changes unmethylated DNA. The DNA is then hybridised to sequences that are either complementary to the original sequence or changed sequence. The hybridisation produces a signal that can be measured.
Methylation-specific multiplex ligation-dependent probe am- plification (MS-MLPA)	In MS-MLPA, the DNA is treated with an enzyme that cleaves unmethylated DNA at specific se- quences, but methylated DNA is protected. PCR to amplify regions of DNA is then performed. Am- plification will only occur if the DNA was not cleaved.
PCR with methylation-sensi- tive high-resolution melting (PCR-HRM)	This technique relies on the changes to DNA caused by sodium bisulfite (i.e. the replacement of cytosine by uracil) leading to it having a lower melting temperature, which is the temperature at which the 2 different DNA strands come apart. Methylated DNA will have a higher melting temperature. A dye that changes fluorescence depending on whether the DNA strands are together or apart can be added.
Co-amplification at lower de- naturation temperature PCR (COLD-PCR)	COLD-PCR relies on the same principle as PCR-HRM. In this case only sequences with low melting temperatures will be amplified. This means that only unmethylated regions will be amplified.
Digestion-based assays	This technique relies on enzymes that cleave unmethylated DNA at specific sequences, but methy- lated DNA is protected.

COLD: co-amplification at lower denaturation temperature; MS-MLPA: methylation-specific multiplex ligation-dependent probe amplification; MSP: methylation-specific polymerase chain reaction; PCR: polymerase chain reaction; PCR-HRM: polymerase chain reaction with high-resolution melting; PSQ: pyrosequencing; qMSP: quantitative methylation-specific polymerase chain reaction.

Table 2. Review question in PICOTS format

Population	People with diagnosed glioblastoma (at any point after diagnosis) who go onto be treated with temozolomide
Index prognostic factors	Tests for MGMT promoter methylation. We considered each method as a separate prognostic fac- tor.
Outcome	Overall survival
Timing	The outcome is to be predicted at any point after the start of treatment.
Setting	To give prognostic information before the start of treatment with temozolomide.

MGMT: O⁶-methylguanine–DNA methyltransferase; PICOTS: Population, Index prognostic factor, Comparator prognostic factor(s), Outcome, Timing, Setting.

Study	ІНС	MSP	PSQ	qMSP	Bead ar- ray	MS-MLPA	PCR-HRM	PCR-mR- NA	Othe
Almuqate 2018	_	_	_	_	_	_	_	_	2
Bady 2012 (E-GBM)	_	_	1	_	2	_	_	_	_
Bady 2012 (M-GBM)	_	1	_	_	1	_	_	-	_
Barault 2015	_	_	1	_	_	_	_	_	1
Barbagallo 2014	_	2	2	_	_	_	_	_	_
Bell 2017	_	_	_	1	_	_	_	_	1
Brigliadori 2016	_	_	2	_	_	_	_	_	_
Chai 2018 (7-site cohort)	_	_	3	_	_	_	_	_	_
Chai 2018 (8-site cohort)	_	_	3	_	_	_	_	-	_
Dahlrot 2018 (NS cohort)	_	_	1	_	_	_	_	_	1
Dahlrot 2018 (RSD cohort)	_	_	1	_	_	_	_	_	1
Dunn 2009	_	_	6	_	_	_	_	_	_
Felsberg 2009	1	1	_	_	_	_	_	1	_
Havik 2012	_	1	9	2	_	_	1	_	_
Hsu 2015	1	1	1	2	_	_	_	_	_
Karayan-Tapon 2010	1	1	6	1	_	_	_	1	_
Kim 2016	_	1	1	_	_	_	_	_	_
Kristensen 2016	1	_	1	_	_	_	_	_	3
Lalezari 2013	1	1	1	_	_	_	_	_	_
Lattanzio 2015	_	2	2	_	_	_	_	_	_

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Table 3. Table of comparisons m	ade (Continued)								
Lechapt-Zalcman 2012	1	1	_	_	_	_	—	—	_
McDonald 2013	_	1	1	_	_	_	_	_	_
Melguizo 2012	1	1	_	_	_	_	_	_	_
Nguyen 2015	_	_	_	2	_	_	_	_	_
Park 2011	_	1	_	_	_	2	_	_	_
Quillien 2016	_	_	12	5	_	_	_	_	_
Quillien 2014 (test)	1	1	32	1	_	_	1	_	_
Quillien 2014 (validation)	_	_	3	_	_	_	_		_
Thon 2017	_	1	_	_	_	_	_		1
Yamashita 2018	_	1	_	-	_	_	5		_
Yang 2012	1	1	_	_	_	_	_		_
Yoshioka 2018	_	_	_	5	_	_	_	_	_

Numbers in cells indicate the number of variants of that technique in the respective study for which we could extract hazard ratios.

IHC: immunohistochemistry; MSP: methylation-specific polymerase chain reaction; PSQ: pyrosequencing; qMSP: methylation-specific polymerase chain reaction; MS-MLPA: methylation-specific multiplex ligation-dependent probe amplification; PCR-HRM: polymerase chain reaction with high-resolution melting; PCR-mRNA: polymerase chain reaction-messenger ribonucleic acid.

Table 4. Impact of adjustment for other prognostic factors

Study ID	Tech- nique	Sample type	CpGs analysed (PCR- based tests)	Thresh- old for methylat- ed	Unadjusted HR (95% CI)	Adjusted HR (95% CI)	Factors adjusted for (if an ad- justed HR reported)	Source of results	Risk of bias (Do- main 6)
Bady 2012 (M-GBM)	Bead array	Frozen	31 and 83	> 0.358	6.46 (2.41 to 17.35)	6.51 (2.42 to 17.54)	Age	IPD	Low

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	MSP	NR	76–80 and 84–86	NR	7.21 (2.37 to 21.99)	7.38 (2.41 to 22.60)	Age	IPD	Low
Dahlrot 2018 (NS cohort)	DIF	FFPE	N/A	< 0.2	1.60 (0.95 to 2.71)	1.50 (0.93 to 2.43)	Age, ECOG performance status and gender	Directly re- ported	Low
Hsu 2015	IHC	FFPE	N/A	< 10%	2.12 (1.32 to 3.42)	1.80 (1.01 to 3.21)	Age, sex, KPS, extent of resec- tion, bevacizumab treatment and IDH1 status	Directly re- ported	Low
	MSP	FFPE	76–80 and 84–87	NR	2.39 (1.42 to 4.02)	2.62 (1.50 to 4.55)	Age, sex, KPS, extent of resec- tion, bevacizumab treatment and IDH1 status	Directly re- ported	Low
	PSQ	FFPE	76–79	> 5%	2.66 (1.49 to 4.76)	2.51 (1.46 to 4.33)	Age, sex, KPS, extent of resec- tion, bevacizumab treatment and IDH1 status	Directly re- ported	Low
	qMSP	FFPE	77–80 and 84–87	> 0.04%	2.75 (1.51 to 5.04)	2.65 (1.47 to 4.76)	Age, sex, KPS, extent of resec- tion, bevacizumab treatment and IDH1 status	Directly re- ported	Low
McDonald 2013	MSP	FFPE	76-80	NR	1.64 (0.95 to 2.83)	1.63 (0.95 to 2.81)	Age	IPD	Low
	PSQ	FFPE	74-78	> 8%	1.96 (1.16 to 3.33)	1.68 (0.99 to 2.84)	Age	Directly re- ported	Low
Thon 2017	MSP	Frozen	76–80 and 84–87	NR	3.33 (1.82 to 6.25)	3.23 (1.72 to 6.25)	Age, RTOG and KPS	Directly re- ported	Low
	Sequenc- ing	Frozen	75–99 (un- clear)	> 50%	3.33 (1.82 to 6.25)	3.23 (1.72 to 6.25)	Age, RTOG and KPS	Directly re- ported	Low
Yamashita 2018	MSP	Frozen	76–80 and 84–87	NR	2.36 (1.62 to 5.05)	1.63 (0.86 to 3.09)	Surgery (gross-total resection vs other) and MGMT status by PCR- HRM	Directly re- ported	High (model includes other MGMT st tus using alternati method)

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Table 4. Impact of adjustment for other prognostic factors (Continued)

	PCR-HRM	Frozen	72-89	> 10%	2.51 (1.63 to 4.83)	2.36 (1.20 to 4.65)	Surgery (gross-total resection vs other) and MGMT status by MSP	Directly re- ported	High (model includes other MGMT sta- tus using alternative method)
Yang 2012	IHC	FFPE	N/A	< 10%	1.07 (0.35 to 3.31)	1.50 (0.37 to 6.04)	Age and extent of resection	IPD	Low
	MSP	FFPE	76–80 and 84–87	NR	1.35 (0.44 to 4.16)	0.99 (0.29 to 3.45)	Age and extent of resection	IPD	Low

CpG: 5'-cytosine-phosphate-guanine-3'; Cl: confidence interval; DIF: double immunofluorescence; ECOG: Eastern Cooperative Oncology Group; FFPE: formalin-fixed paraffin embedded; HR: hazard ratios; IDH: isocitrate dehydrogenase; IHC: immunohistochemistry; IPD: individual participant data; KPS: Karnofsky performance status; MGMT: O⁶methylguanine–DNA methyltransferase; MSP: methylation-specific polymerase chain reaction; N/A: not applicable; NR: not reported; PCR: polymerase chain reaction; PCR-HRM: polymerase chain reaction with high-resolution melting; PSQ: pyrosequencing; qMSP: quantitative methylation-specific polymerase chain reaction; RTOG: Radiation Therapy Oncology Group prognostic factor class.

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Table 5. Details of articles describing a single method

Author (year)	Country of popula- tion/cohort	Length of follow-up (median in months (range))	Number of partici- pants	Tumour type	IDH muta- tion status (% of WT)	Technique used to as- sess MGMT methyla- tion status
Abhinav 2013	UK	NR	19	GBM	NR	MSP
Adeberg 2015	Germany	NR	32	GBM	NR	MSP
Ahmed 2015	US	11.4	214	GBM	NR	mRNA lev- els
Alonso 2017	Spain	NR	63	GBM: 97.2%; gliosarcoma: 2.8%	88.7 (data from 63/71 people)	MSP
Appin 2013	US	NR	236	GBM: prima- ry: 89% (GBM with oligoden- droglioma: 13.3%); sec- ondary: 11%	GBM: 91.4 (data from 116/208 peo- ple); GBM- O: 65 (data from 20/28 people)	MSP
Ardon 2012	Belgium	25.0 (10.5– 42.2)	77	GBM (primary)	NR	MSP
Arita 2016	Japan	NDR	193	GBM	100	PSQ
Badruddoja 2017	US	NR	30	GBM	NR	qMSP
Balana 2016	Spain	NR	93	GBM	NR	MSP
Balana 2017	Spain	17.0 (10.7– 24.5)	256	GBM	94.5 (data from 162/256 people)	MSP
Blumenthal 2017	US	NR	1395	GBM	NR	qMSP
Boots-Sprenger 2013	The Netherlands	NR	333	GBM	84 (data from 226/333 people)	MS-MLPA
Brandes 2008	Italy	18.93 (6.6– 62)	208	GBM	NR	MSP
Brandes 2009	Italy	NR	37	GBM	NR	MSP
Brandes 2010	Italy	NR	44	GBM	NR	MSP
Brandes 2014	Italy	NR	116	GBM	NR	MSP
Brandes 2017	Italy	NR	108	GBM	NR	MSP

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 78



Table 5. Details of articles describing a single method (Continued)

Brennan 2013	US	NR	332	GBM	93.4 (data from 423/543 people)	Bead array
Burford 2013	UK	NR	NDR	GBM	NDR	MSP
Burger 2017	Germany	NR	32	GBM	NR	MSP
Butowski 2011	US	NR	66	GBM and gliosar- coma	NR	MSP
Capellades 2018	Spain	NR	292	GBM	96.6	MSP
Chakhoyan 2018	US	NR	23	GBM	NR	MSP
Chen 2015	China	12.8 (4.0– 37.7)	78	GBM	NR	IHC
Chen 2016	China	NR	300	GBM	85.7	Bead array
Cheng 2015	China, US	NR	285 (CGGA: 55; TCGA: 235)	GBM	NR	Bead ar- ray (TCGA); PSQ (CGGA)
Chinot 2007	France	6 (0.9–19)	29	GBM	NR	IHC
Choi 2016	South Korea	NR	112 (train- ing cohort: 74; test co- hort 38)	GBM	NR	MSP
Clarke 2009	US	18.8	85	GBM	NR	MSP
Coburger 2017	Germany	40 (37–43)	170	GBM	NR	MSP
Colman 2010	US	NR	101	GBM	NR	qMSP
Combs 2011	Germany	NR	160	GBM (primary)	97.1 (data from 140/160 people)	MSP
Cominelli 2015	Italy	NR	70	GBM	95.7	MSP
Costa 2010	Portugal	NR	90	GBM (primary)	NR	MSP
Criniere 2007	France	57.2	77	GBM	NR	MSP
Dahlrot 2017	Denmark	NR	226	GBM (primary)	NR	PSQ
Das 2011	India	NR	6	GBM	NR	MSP
Etcheverry 2014	France	15.5	399	GBM (primary)	91	PSQ
Felsberg 2011	Germany	48.6	64	GBM	NR	MSP
Fiano 2014	Italy	NR	32	GBM	NR	MSP

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 79



Table 5. Details of articles describing a single method (Continued)

Fontana 2016	Italy	NDR	128	GBM	NDR	PSQ
Franceschi 2016	Italy	NR	21	GBM	NR	MSP
Franceschi 2018	Italy	NR	169	GBM	NR	MSP
Galldiks 2015	Germany	NR	21	GBM	NR	MSP
Gallego Perez-Larraya 2011	France	NR	31	GBM	NR	qMSP
Gilbert 2013	Canada, European multicentres, US	NR	762	GBM	NR	qMSP
Gilbert 2014	Canada, European multicentres, US	NR	637	GBM	NR	qMSP
Gittleman 2017	US	NR	799	GBM	NR	qMSP
Glas 2009	Germany	41.5	23	GBM	NR	MSP
Gorlia 2008	Belgium, Cana- da, Italy, Germany, Switzerland, the Netherlands	NR	287	GBM	NR	MSP
Gramatzki 2016	Switzerland	9	108	GBM	100	MSP
Gutenberg 2013a	Germany	NR	17	GBM (primary)	NR	MSP
Gutenberg 2013b	Germany	16.2 (1.4– 54.1)	191	GBM (primary)	NR	MSP
Ha 2013	South Korea	NR	10	GBM	75	qMSP
Haemmig 2014	Switzerland	NR	60	GBM	85	qMSP
Han 2014	US	NR	28	GBM	NR	MSP
Han 2015a	China	13.7 (1–43)	152	GBM	NR	MSP
Han 2015b	China	13.7 (1–43)	79	GBM	93.5 (data from 214 people)	MS-MLPA
Happold 2018	European multicen- tres	29 (25–35; CENTRIC cohort)	797	GBM	NR	qMSP
Hayes 2015	US	14.1	219	GBM	94.5 (data from 475 people)	Bead array
Hegi 2004	Switzerland	NR	38	GBM	NR	MSP
Hegi 2005	Canada, European multicentres	NR	106	GBM	NR	MSP

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 80



Table 5. Details of articles describing a single method (Continued)

Herrlinger 2006	Germany	NR	31	GBM	NR	MSP
Herrlinger 2009	Germany	41.5	31	GBM	NR	MSP
Hervouet 2009	France	NR	53	GBM	NR	MSP
Hobbs 2012	US	19.1 (8.1– 74.6; sur- vivors only)	312	GBM	NR	qMSP
Huang 2017	US	NR	301	GBM	NR	mRNA lev- els
Hudson 2018	Australia	NR	16	GBM	NR	PSQ
Inoges 2017	Spain	NR	31	GBM	NR	MSP
Ishida 2015	Japan	NR	46	GBM	NR	IHC
Ishikawa 2014	Japan	19.6 (7.3– 48.7)	23	GBM	75 (data from the whole co- hort)	IHC
lus 2018	Italy	NR	116	GBM	NR	PSQ
Iwadate 2017	Japan	NR	70	GBM	92	IHC
Jan 2018	Taiwan	NR	NDR	GBM	NDR	MSP
Karim 2012	Egypt	NR	34	GBM	NR	MSP
Kessler 2018	Germany, US	NR	404 (Hei- delberg co- hort: 143; TCGA: 261)	GBM	100	Bead array
Kim 2012	South Korea	22 (3-88)	93	GBM	NR	MSP
Kim 2017	South Korea	16.3 (0.3– 105.1)	750	GBM	NR	MSP
Kim 2018	South Korea	NR	93	GBM	86.8 (data from 91 peo- ple)	qMSP
Klitkou 2014a	Denmark	NR	173	GBM (primary)	NR	IHC
Klitkou 2014b	Denmark	NR	173	GBM (primary)	NR	IHC
Klitkou 2014c	Denmark	NR	173	GBM (primary)	NR	IHC
Kong 2011	South Korea	16.5 (6.2– 48)	90	GBM	NR	MSP
Kreth 2013	Germany	NR	222	GBM	NR	MSP

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)



Table 5. Details of articles describing a single method (Continued)

Lakomy 2011	Czech Republic	NR	38	GBM	NR	PCR-HRM
Laxton 2013	UK	NR	288	GBM	95.3 (data from 107/288 people)	MSP
Lee 2013	South Korea	15	36	GBM	NR	qMSP
Lee 2017	South Korea	NR	65	GBM	80	MSP
Li 2016a	China	NR	145	GBM	83.4	MSP
Li 2016b	China	NR	50	GBM (primary)	85.9 (data from 50/78 people)	PSQ
Lombardi 2015	Italy	NR	151	GBM	94 (data from 100/237 people)	MSP and PSQ
Lombardi 2017	Italy	NR	128	GBM	NR	PSQ
Ma 2016	China	NR	56	GBM (primary)	NR	MSP
Majewska 2017	United Kingdom	NR	99	GBM	NR	PSQ
Malmström 2012	Austria, Denmark, France, Norway, Sweden, Switzer- land, Turkey	NR	72	GBM	99.7 (data from 291 people)	qMSP
Malmström 2017	Denmark, Finland, Norway, Sweden	20	78	GBM	96.10%	PSQ
Martini 2008	Italy	NR	46	GBM	NR	MSP
McDonald 2015	Australia	NR	33	GBM	93.90%	PSQ
Metellus 2011	France	18.9	61	GBM	NR	qMSP
Meyronet 2017	Austria, Denmark, France, Norway, Sweden, Switzer- land, Turkey	NR	6	GBM	100	PSQ
Michaelsen 2013	Denmark	60 (23–92)	163	GBM	NR	IHC
Michaelsen 2018	Denmark	NR	415	GBM	100	mRNA lev- els
Minniti 2011a	Italy	NR	36	GBM (recurrent)	NR	MSP
Minniti 2011b	Italy	NR	83	GBM	NR	MSP
Minniti 2015	Italy	24.0 (stan- dard RT + TMZ group);	243	GBM	NR	MSP

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 82



Table 5. Details of articles describing a single method (Continued)

		22.5 (short course RT + TMZ group)				
Miyazaki 2014	Japan	NR	117	GBM: 83.8%; GBM with oligo- dendroglioma: 16.2%	93.4	IHC
Montano 2011	Italy	NR	73	GBM (primary)	NR	MSP
Morandi 2010	Italy	NR	159	GBM	NR	qMSP
Motomura 2011	Japan	16.7 (3.4– 46.7)	68	GBM (primary)	94.1	PSQ
Mur 2015	Spain	NR	68	GBM	70.6 (un- known in 25%)	Bead array
Nabors 2012	US	NR	69	GBM	NR	qMSP
Nagane 2007	Japan	7.1 (2.4– 16.7)	19	GBM (recurrent)	NR	Western blot analy- sis
Ohka 2011	Japan	NR	51	GBM (primary)	94	PSQ
Ohno 2013	Japan	NR	85	GBM	NR	PSQ
Ohno 2016	Japan	NR	112	GBM	92	PSQ
Omuro 2014	US	42	40	GBM	100	qMSP
Pallini 2008	Italy	NR	44	GBM	NR	MSP
Pambuku 2016	Italy	NR	128	GBM	NR	PSQ
Park 2013	South Korea	NR	75	GBM	NR	MSP
Pei 2013	China	NR	54	GBM	NR	MSP
Picart 2018	France	NR	14	GBM (cerebellar)	100	PSQ
Poulsen 2017	Denmark	14	146	GBM	98	IHC
Prados 2009	US	33.7	65	GBM and gliosar- coma	NR	MSP
Purkait 2016	India	NR	114	GBM	93.3	MSP
Qi 2012	China	NR	86	GBM (secondary)	26.6 (data from 79 peo- ple)	MSP
Rankeillor 2014	UK	NR	29	GBM (primary)	NR	MS-MLPA

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Table 5. Details of articles describing a single method (Continued)

Rapkins 2015	Australia, US	NR	319 (AGOG cohort: 160; UCLA co- hort: 159)	GBM	NR	MSP (UCLA); PSQ (AGOG)
Rapp 2013	Germany	NR	85	GBM (primary)	NR	MSP
Reifenberger 2012	Germany	29	104	GBM	NR	MSP
Roh 2017	South Korea	20.8	252	GBM: 84.5%; gi- ant-cells GBM: 4.0%; GBM with oligoden- droglioma: 11.5%	93.4 (data from 106/252 people)	MSP
Romano 2013	Italy	NR	47	GBM	NR	MSP
Rosati 2013	Italy	11.5 (1.5– 58)	83	GBM: 95.2%; gliosarcoma: 2.4%; GBM with oligodendroglia: 2.4%	97.6	MSP
Rosenschold 2019	Denmark	NR	412	GBM (primary)	NR	IHC
Rubio Fernandez 2014	Spain	NR	65	GBM	NR	MSP
Sadones 2009	Belgium	NR	32	GBM (recurrent)	NR	qMSP
Saito 2017a	Japan	NR	53	GBM (supraten- torial)	49 (unknown in 49%)	IHC
Saito 2017b	Japan	NR	36	GBM (supraten- torial)	NR	IHC
Saito 2018a	Japan	NR	102	GBM (supraten- torial)	75.5	IHC
Saito 2018b	Japan	NR	50	GBM (supraten- torial)	100	IHC
Saito 2018c	Japan	NR	50	GBM (supraten- torial)	NR	IHC
Salvati 2012	Italy	NR	105	GBM (primary supratentorial)	NR	MSP
Sana 2014	Czech Republic	NR	58	GBM (primary)	NR	PCR-HRM
Saraiva-Esperon 2014	Spain	NR	25	GBM	NR	MSP
Sasaki 2018	Japan	NR	101	GBM: 99%; gliosarcoma: 1%	99	qMSP
Schaich 2009	Germany	NR	64	GBM (supraten- torial)	NR	MSP

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Table 5. Details of articles describing a single method (Continued)

Schiffgens 2016	Germany, Italy	NR	225 (Han- nover co- hort 1: 120); Bologna cohort: 105)	GBM (primary)	91.4 (da- ta from the Bologna co- hort)	MSP
Schulze Heuling 2017	Germany, US	NR	275	GBM	100	Bead array
Shu 2018	China	NR	265	GBM (primary)	100	PSQ
Sijben 2008	Canada	NR	16	GBM (supraten- torial)	NR	MSP
Singh 2012	India	13.15 (1.5– 46)	16	Gliosarcoma	NR	MSP
Soike 2018	US	NR	74	GBM	89.2	IHC
Stetson 2016	US	NR	203 (train- ing cohort: 102; valida- tion cohort: 101)	GBM	Training cohort: 75 (unknown: 21%); vali- dation co- hort: 77 (un- known: 17%)	Bead array
Stummer 2012	Germany	24	143	GBM	NR	MSP
Stupp 2009	Canada, European multicentres	61 (0.36– 79)	287	GBM	NR	MSP
Stupp 2010	Germany, Switzer- land	34	52	GBM	NR	MSP
Suchorska 2015	Germany	NR	79	GBM (supraten- torial)	NR	MSP
Tanaka 2014	Japan	NR	45	GBM	NR	mRNA lev- els
Thon 2011	Germany	11 (5–33)	56	GBM	NR	MSP
Tini 2015	Italy	NR	144	GBM	NR	MSP
Tini 2016	Italy	12 (6–84)	169	GBM	100	MSP
Tini 2017	Italy	NR	222	GBM	NR	MSP
Toms 2018	Multicentre (North America, Europe, South Korea, Israel)	NR	466	GBM (supraten- torial)	NR	MSP
Trabelsi 2016	Tunisia	NR	20	GBM	NR	MS-MLPA
Urbschat 2017	Germany	NR	72	GBM	NR	MSP

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Table 5. Details of articles describing a single method (Continued)

van Dijken 2019	China, Taiwan, the Netherland, UK	NR	50	GBM	84 (missing in 8%)	PSQ
Villani 2015	Italy	12 (3–27)	51	GBM (primary)	NR	PSQ
Wang 2014	China	47 (6–65)	78	GBM	52.6	MSP
Wang 2015a	China	NR	216	GBM	48.1	MSP
Wang 2015b	Multicentre (North America and Eu- rope)	31.9 (0.2– 53.3)	831	GBM	NR	qMSP
Wang 2016	China, US	NR	84 (CGGA: 21; TCGA: 63)	GBM	NR	Bead ar- ray (TCGA); PSQ (CGGA
Watanabe 2011	Japan	15.4	41	GBM	NR	IHC
Wee 2017	South Korea	20.5	340	GBM	93.8	MSP
Wee 2018	Japan, South Korea	18.4	692	GBM	92.3	MSP
Wei 2017	Taiwan	NR	25	GBM	NR	PSQ
Weller 2015	Austria, Germany, Switzerland	NR	105	GBM (recurrent)	NR	MSP
Weller 2017b	Multicentre (165 hospitals in 22 countries)	NR	745	GBM (EGFRvIII- expressing)	NR	MSP
Weller 2009	Germany	29.4 (data from the whole co- hort)	189	GBM: 96.7; gi- ant cells GBM: 2.6%; gliosarco- ma: 0.75	94.4 (da- ta from the whole co- hort)	MSP
Westphal 2015	Germany	NR	66	GBM	NR	PSQ
Wu 2018	US	NR	285	GBM	NR	Bead array
Yan 2017	UK	NR	31	GBM	90.3	PSQ
Yang 2015	China	NR	229	GBM	76.3 (data from 274 people; un- known in 3.3%)	PSQ
Yin 2017	France	NR	106	GBM	NR	Bead array
Yin 2018	France	53 (8–113; Rennes and Angers datasets)	129	GBM	NR	Bead array
You 2013	Taiwan	NR	32	GBM	NR	qMSP

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Table 5. Details of articles describing a single method (Continued)

Younis 2016Egypt13 (9-27)73GBMNRIHCYuan 2017aChinaNR84GBMNRPSQYuan 2017bChinaNR48GBMNRPSQYue 2014China17.5 (3-77)62GBMNRIHCZhang 2014ChinaNR80GBMNRPCR-HRMZunarelli 2011Italy10.946GBM (primary)NRMSP		• •					
Yuan 2017bChinaNR48GBMNRPSQYue 2014China17.5 (3-77)62GBMNRIHCZhang 2014ChinaNR80GBMNRPCR-HRM	Younis 2016	Egypt	13 (9–27)	73	GBM	NR	IHC
Yue 2014 China 17.5 (3-77) 62 GBM NR IHC Zhang 2014 China NR 80 GBM NR PCR-HRM	Yuan 2017a	China	NR	84	GBM	NR	PSQ
Zhang 2014 China NR 80 GBM NR PCR-HRM	Yuan 2017b	China	NR	48	GBM	NR	PSQ
	Yue 2014	China	17.5 (3–77)	62	GBM	NR	IHC
Zunarelli 2011 Italy 10.9 46 GBM (primary) NR MSP	Zhang 2014	China	NR	80	GBM	NR	PCR-HRM
	Zunarelli 2011	Italy	10.9	46	GBM (primary)	NR	MSP

EGFRvIII: epidermal growth factor receptor variant 3; GBM: glioblastoma; IDH: isocitrate dehydrogenase; IHC: immunohistochemistry; MGMT: O⁶-methylguanine–DNA methyltransferase; mRNA: messenger ribonucleic acid; MS-MLPA: methylation-specific multiplex ligationdependent probe amplification; MSP: methylation-specific polymerase chain reaction; NDR: not directly reported; NR: not reported; PCR-HRM: polymerase chain reaction with high-resolution melting; PSQ: pyrosequencing; qMSP: quantitative methylation-specific polymerase chain reaction; RT: radiotherapy; TMZ: temozolomide; WT: wild-type.

APPENDICES

Appendix 1. Database search strategies

Date of search: 3 December 2018

- Ovid MEDLINE (1946 to 4 December 2018), 1500 records.
- PubMed NOT MEDLINE (4 December 2018), 101 records.
- Ovid Embase (1980 to 2018, week 49), 2983 records.
- BIOSIS (1969 to 3 December 2018), 790 records.
- Web of Science Conference Proceedings Citation Index (CPCI-S) (1900 to 3 December 2018), 120 records.

Total: 5494 records

Duplicates removed: 2137 records

Records to screen: 3357

Ovid MEDLINE(R) and Epub Ahead of Print, In-Process & Other Non-Indexed Citations and Daily <1946 to 4 December 2018>

Search strategy:

1 glioma/ or astrocytoma/ or glioblastoma/ (65202)

2 (glioblastom* or GBM or astrocytom* or gliosarcom*).mp. (55603)

3 1 or 2 (81995)

4 "O(6)-Methylguanine-DNA Methyltransferase"/ (2192)

5 ((methylguanin* or methyl guanin* or alkylguanin* or alkyl guanin*) adj5 (methyltransferas* or methyl transferas* or alkyltransferas* or alkyl transferas* or transmethylas* or trans methylas*)).mp. (3857)

6 (methyl* DNA protein cystein* adj (methyltransferas* or methyl transferas*)).mp. (3)

7 (AGT or MGMT or AGAT).ti,ab,kf,ot. (6403)

8 or/4-7 (7811)

9 exp Prognosis/ (1466623)

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10 (prognos* or predict*).mp. (2042311)

- 11 exp mortality/ (350654)
- 12 survival/ (4532)
- 13 survival rate/ (158119)
- 14 exp survival analysis/ (264438)
- 15 (mortality or death* or surviv*).mp. (2412078)
- 16 Follow-Up Studies/ (602886)
- 17 ((followup or follow-up) adj (study or studies)).ti,ab,kf. (48020)
- 18 or/9-17 (4920356)
- 19 3 and 8 and 18 (1515)
- 20 exp animals/ not humans.sh. (4519948)
- 21 19 not 20 (1500)
- *****

Ovid Embase <1980 to 2018 week 49>

Search strategy:

1 glioma/ or astrocytoma/ or glioblastoma/ (104447)

2 (glioblastom* or GBM or astrocytom* or gliosarcom*).mp. (88786)

3 1 or 2 (119931)

4 methylated DNA protein cysteine methyltransferase/ (5065)

5 (methyl* DNA protein cystein* adj (methyltransferas* or methyl transferas*)).ti,ab,kw,ot. (3)

6 ((methylguanin* or methyl guanin* or alkylguanin* or alkyl guanin*) adj5 (methyltransferas* or methyl transferas* or alkyltransferas* or alkyl transferas* or transmethylas* or trans methylas*)).mp. (4545)

7 (AGT or MGMT or AGAT).ti,ab,kw,ot. (10397)

8 or/4-7 (13071)

- 9 prognosis/ or cancer prognosis/ (629552)
- 10 (prognos* or predict*).mp. (2614430)
- 11 exp mortality/ (931698)
- 12 exp survival/ (972881)
- 13 survival analysis/ (12408)
- 14 (mortality or death? or surviv*).ti,ab,kw,ot. (2769350)
- 15 or/9-14 (4989016)
- 16 Methylation/ or DNA methylation/ (107678)
- 17 methylat*.ti,ab,kw,ot. (123232)



18 ((amount? or express* or level? or activ* or status) adj5 (protein? or AGT or MGMT or AGAT or ((methylguanin* or methyl guanin* or alkylguanin* or alkylguanin* or alkylguanin*) adj5 (methyltransferas* or methyl transferas* or alkyltransferas* or alkyl transferas* or transmethylas* or transmethylas*)))).ti,ab,kw,ot. (962065)

19 or/16-18 (1093057)

20 3 and 8 and 15 and 19 (3054)

21 ((animal or nonhuman) not human).de. (5171408)

22 (cell line or cell culture).hw. not ((human or adult).sh. or patient.hw.) (344719)

23 (20 not (21 or 22)) (2983)

BIOSIS Citation Index (3 December 2018)

[Search-1: ((tumour 'near' enzyme) and prognosis)]

#1 ((TS=((glioblastoma* OR GBM* OR astrocytom*) NEAR (methylguanin* OR "methyl guanin*" OR alkylguanin* OR "alkyl guanin*" OR AGT OR MGMT OR AGAT)) OR TS=((gliosarcom*) AND (methylguanin* OR "methyl guanin*" OR alkylguanin* OR "alkyl guanin*" OR AGT OR MGMT OR AGAT))) <u>AND</u> (TS=(prognos* or predict* or mortalit* or death* or surviv*)))n=437

[Search-2: ((prognosis 'near' enzyme) and tumour)]

#2 TS=((prognos* OR predict* OR mortalit* OR death* OR surviv*) NEAR (methylguanin* OR "methyl guanin*" OR alkylguanin* OR "alkyl guanin*" OR AGT OR MGMT OR AGAT)) <u>AND</u> TS=(glioblastom* OR GBM* OR astrocytom* OR gliosarcom*) n=425

[Search-3: ((prognosis 'near' tumour) and enzyme)]

#3 ((TS=((prognos OR predict* OR mortalit* OR death* OR surviv*) NEAR (glioblastom* OR GBM* OR astrocytom*)) OR TS=((prognos OR predict* OR mortalit* OR death* OR surviv*) AND gliosarcom*)) <u>AND</u> (TS=("O(6)-Methylguanine-DNA Methyltransferase" OR "O-6-Methylguanine-DNA Methyltransferase" or "methylated DNA protein cysteine methyltransferase" OR AGT or MGMT or AGAT) OR TS=((methylguanin* or "methyl guanin*") or alkylguanin* or "alkyl guanin*") NEAR (methyltransferas* or "methyl transferas*" or alkyltransferas* or "transmethylas* or "transmethylas*"))) n=413

[Search-4: prognosis AND tumour AND enzyme AND methylation/expression]

#4 (TS=(glioblastom* or GBM or astrocytom* or gliosarcom*) AND (TS=("O(6)-Methylguanine-DNA Methyltransferase" or "O-6-Methylguanine-DNA Methyltransferase" or "methylated DNA protein cysteine methyltransferase" or AGT or MGMT or AGAT) OR TS=((methylguanin* or "methyl guanin*" or alkylguanin* or "alkyl guanin*") NEAR (methyltransferas* or "methyl transferas*" or alkyltransferas* or "alkyl transferas*" or transmethylas* or "trans methylas*"))) AND (TS=(prognos* or predict* or mortality or death or deaths or surviv*)) AND (TS= methylat* OR TS=((amount or amounts or express* or level or levels or activ* or status) NEAR (protein* or AGT or MGMT or AGAT or methylguanin* or "methyl guanin*" or alkylguanin* or "alkyl guanin*")))) n=722

#5 (#4 or #3 or #2 or #1) n=790

[N.B. Gliosarcoma is a much rarer tumour, so the proximity operator was not used in this context, in search lines 1 and 3)]

Conference Proceedings Citation Index- Science (CPCI-S) (1990 to 3 December 2018)

#1 TS=((glioblastom* or GBM or astrocytom* or gliosarcom*) and (prognos* or predict* or mortality or death or deaths or surviv*)) n=2200

#2 TS=("O(6)-Methylguanine-DNA Methyltransferase" or "O-6-Methylguanine-DNA Methyltransferase" or "methylated DNA protein cysteine methyltransferase" or AGT or AGAT) n=1042

#3 TS=((methylguanin* or "methyl guanin*" or alkylguanin* or "alkyl guanin*") AND (methyltransferas* or "methyl transferas*" or alkyltransferas* or "alkyl transferas*" or transmethylas* or "trans methylas*")) n=277

#4 (#3 OR #2) n=1135

#5 (#4 and #1) n=120

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PubMed 4 December 2018

#12 Search (#10 AND #11) (101)

#11 Search pubmednotmedline[sb] (2531482)

#10 Search (#3 AND #9) (1855)

#9 Search (#4 OR #5 OR #6 OR #7 OR #8) (11388)

#8 Search (AGT[Title/Abstract] OR MGMT[Title/Abstract] OR AGAT[Title/Abstract]) (6400)

#7 (("methylated DNA protein cysteine methyltransferas*" or "methylated DNA protein cysteine methyl transferas*" or "methylguanine deoxyribonucleic acid methyltransferas*" or "methylguanine deoxyribonucleic acid methyl transferas*" or "methyl guanine deoxyribonucleic acid methyl transferas*" or "methylguanine deoxyribonucleic acid methyl transferas*" or "methyl guanine deoxyribonucleic acid methyl transferas*" or "methyl guanine deoxyribonucleic acid methyl transferas*" or "methylguanin* dna protein methyltransferas*" or "methyl guanin* dna protein methyl transferas*" or "methyl guanin* dna protein methyl guanin* dna protein methyl guanin* dna protein methyl guanin* dna protein methyl guanin* dna protein methyl

#6 (("methylguanin* DNA methyltransferas*" OR "methylguanin* DNA methyl transferas*" OR "methylguanin* DNA alkyltransferas*" OR "methylguanin* DNA alkyltransferas*" OR "methyl guanin* DNA methyl guanin* DNA methyl transferas*" OR "methyl guanin* DNA alkyltransferas*" OR "methyl guanin* DNA alkyltransferas*" OR "methyl guanin* DNA alkyltransferas*" OR "alkylguanin* DNA methyl transferas*" OR "alkylguanin* DNA alkyltransferas*" OR "alkylguanin* DNA methyl transferas*" OR "alkylguanin* DNA alkyltransferas*" OR "alkyl

#5 Search (("methylguanin* methyltransferas*" OR "methylguanin* methyl transferas*" OR "methylguanin* alkyltransferas*" OR "methyl guanin* methyl transferas*" OR "methyl guanin* alkyltransferas*" OR "methyl guanin* methyl transferas*" OR "methyl guanin* alkyltransferas*" OR "methyl guanin* methyl transferas*" OR "methyl guanin* alkyltransferas*" OR "alkylguanin* methyl transferas*" OR "alkylguanin* alkyltransferas*" OR "alkylguanin* alkyl transferas*" OR "alkylguanin* methyl guanin* methyl transferas*" OR "alkylguanin* methyl transferas*" OR "alkylguanin* alkyl transferas*" OR "alkylguanin* methyl guanin* methyl transferas*" OR "alkylguanin* methyl guanin* methyl transferas*" OR "alkylguanin* alkyl transferas*" OR "alkylguanin* methyl guanin* methyl guanin* methyl transferas*" OR "alkylguanin* methyl guanin* methyl guan

#4 Search "O(6)-Methylguanine-DNA Methyltransferase"[Mesh:NoExp] (2192)

#3 Search (#1 OR #2) (81976)

#2 Search (glioblastom* OR GBM OR astrocytom* OR gliosarcom*) (55573)

#1 ("Glioma"[Mesh:NoExp] OR "Astrocytoma"[Mesh:NoExp] OR "Glioblastoma"[Mesh:NoExp]) (65225)

Appendix 2. Risk of bias and applicability assessment

Bespoke tool to assess risk of bias and applicability of prognostic factor studies. SQ: signalling question.

Domain 1: participant	selection
Risk of bias	SQ1.1: was a consecutive or random sample of people enrolled?
	SQ1.2: was a case-control or cross-sectional design avoided?
	SQ1.3: did the study avoid inappropriate exclusions?
Applicability	Are there concerns that the included participants and setting do not match the review question?
Domain 2: subsequent	treatment
Risk of bias	SQ2.1: did treatment vary across participants? (or "Was treatment either standardised or ran- domised?")
Applicability	Are there concerns that treatments received do not match the review question?

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(Continued)

Domain 3: outcome measu	rement				
Risk of bias	SQ3.1: was the method of outcome measurement used adequately valid and reliable?				
	SQ3.2: was the method and setting of outcome measurement the same for all study participants?				
	SQ3.3: was the outcome objective or assessed without knowledge of the prognostic factor?				
	SQ3.4: do the prognostic factors investigated form part of the outcome?				
Applicability	Are there concerns that outcome does not match the question or that follow-up was not of suffi- cient duration, or both?				
Domain 4: prognostic facto	or measurement				
Risk of bias	SQ4.1: was the method and setting of measurement of the prognostic factor the same for all partic ipants?				
	SQ4.2: was the prognostic factor objective or measured without knowledge of the outcome or risk of the outcome?				
	SQ4.3: if a threshold was used, was it prespecified?				
Applicability	Are there concerns that prognostic factor, the way that it was measured, or the way that it was i terpreted, differ from the review question?				
Domain 5: study attrition					
Risk of bias	SQ5.1: were all participants included in the analysis?				
	If no to SQ5.1: SQ5.2: were there important differences between participants who completed the study/were included in the analysis and those who were not?				
Domain 6: adjustment for	other potential prognostic factors (where relevant)				
Risk of bias	SQ6.1: were other potential prognostic factors measured adequately and reliably and in a similar manner for all participants, and is the method of adding them to the model appropriate?				
Applicability	Did the prognostic factors adjusted for match the review question?				
Domain 7: selective report	ing				
Risk of bias	SQ7.1: is the reported estimate likely to be selected on the basis of the results from: multiple out- come measurements, multiple analyses of the prognostic factor-outcome relationship, from differ ent subgroups, or a combination of these?				

Appendix 3. Domains to be considered when judging the strength of the body of evidence

We considered the following domains when we assessed the strength of the body of evidence, based on the GRADE approach (Guyatt 2008).

Domain

Explanation

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 91 people with glioblastoma treated with temozolomide (Review)



(Continued)	
Risk of bias	Based on results of 'Risk of bias' assessments, we downgraded confidence in the evidence base if most evidence was from studies that we judged at high risk of bias.
Indirectness	We downgraded confidence in the evidence base if we had concerns that the study sample, the prognostic factor, the outcome, other factors in the models in the primary studies or a combination of these did not reflect the review question.
Inconsistency	We downgraded confidence in the evidence base if there was unexplained heterogeneity or vari- ability in results across studies.
Imprecision	We downgraded confidence in the evidence base if the estimate of the effect size from a meta- analysis was not precise or, if no meta-analysis was performed, if the estimate of the size of effect from individual studies was not precise.
Publication bias	Studies showing no association are likely to be unpublished, unless part of a larger study that specifically aimed to compare tests. We downgraded our confidence in the evidence base if we had reason to suspect publication bias from our assessments of reporting bias.
Size of effect	We upgraded our confidence in the evidence base if the size of effect was moderate or large. If a meta-analysis is not possible, we upgraded if the size of effect was moderate or large for most included studies.

Appendix 4. Identification of studies making different comparisons among the techniques

Comparison numbers are for [row-defining technique] versus [column-defining technique], for example Kristensen 2016 (top right cell) compares one version of IHC versus three 'other' techniques.

 Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in
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 people with glioblastoma treated with temozolomide (Review)
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	IHC	MSP	PSQ	qMSP	Bead ar- ray	MS-MLPA	PCR-HRM	PCR-mR- NA	Other
ΙΗС	None	Felsberg 2009 (1 vs 1) Hsu 2015 (1 vs 2) Karayan- Tapon 2010 (1 vs 1) Lalezari 2013 (1 vs 1) Lechapt-Zal- cman 2012 (1 vs 1)	Hsu 2017 (1 vs 1) Karayan-Tapon 2010 (1 vs 6) Kristensen 2016 (1 vs 1) Lalezari 2013 (1 vs 1) Quillien 2014 (test) (1 vs 32)	Hsu 2017 (1 vs 2) Karayan-Tapon 2010 (1 vs 1) Quillien 2014 (test) (1 vs 1)	None	None	Quillien 2014 (test) (1 vs 1)	Felsberg 2009 (1 vs 1) Karayan- Tapon 2010 (1 vs 1)	Kristensen 2016 (1 vs 3
		Melguizo 2012 (1 vs 1) Quillien 2014 (test) (1 vs 1) Yang 2012 (1 vs 1)							
MSP	_	Barbagallo 2014 (2) Hsu 2017 (2) Lattanzio 2015 (2)	Barbagallo 2014 (2 vs 2) Havik 2012 (1 vs 9) Hsu 2017 (2 vs 1) Karayan-Tapon 2010 (1 vs 6) Kim 2016 (1 vs 1) Lalezari 2013 (1 vs 1) Lattanzio 2015 (1 vs 2) McDonald 2013 (1 vs 1); Quillien 2014 (test) (1 vs 32)	Havik 2012 (1 vs 2) Hsu 2017 (2 vs 2) Karayan-Tapon 2010 (1 vs 1) Quillien 2014 (test) (1 vs 1)	Bady 2012 (M-GBM) (1 vs 1)	Park 2011 (1 vs 2)	Havik 2012 (1 vs 1) Quillien 2014 (test) (1 vs 1) Yamashita 2018 (1 vs 5)	Felsberg 2009 (1 vs 1) Karayan- Tapon 2010 (1 vs 1)	Thon 2017 (1 vs 1)

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(Continued)									
PSQ	-	—	Barbagallo 2014 (2)	Havik 2012 (9 vs 2)	Bady 2012 (E-GBM) (1	None	Havik 2012 (9 vs 1)	Karayan- Tapon	Barault 201 (1 vs 1)
			Brigliadori 2016 (2) Chai 2018 (7-site cohort) (3)	Hsu 2017 (1 vs 2)	vs 2)		Quillien	2010 (6 vs 1)	Dahlrot
			Chai 2018 (8-site cohort) (3)	Karayan-Tapon 2010 (6 vs 1)			2014 (test) (32 vs 1)		2018 (NS Cohort) (1 v 1)
			Dunn 2009 (6)	Quillien 2014) Dahlrot
			Havik 2012 (9)	(test) (32 vs 1)				2018 (RSD Cohort) (1 v	
			Karayan-Tapon 2010 (6)	Quillien 2016 (12 vs 5)					1)
			Lattanzio 2015 (2)						Kristensen 2016 (1 vs 3
			Quillien 2014 (test) (32)						(
			Quillien 2014 (validation) (3)						
			Quillien 2016 (12)						
qMSP	_	—	_	Havik 2012 (2)	None	None		Karayan- Tapon	Bell 2017 (1 vs 1)
				Hsu 2017 (2)			Quillien	2010 (1 vs 1)	,
				Nguyen 2015 (2)			2014 (test) (1 vs 1)	1)	
				Quillien 2016 (5)		(1 03 1)	(1 00 1)		
				Yoshioka 2018 (5)					
Bead ar- ray	_	_	_	_	Bady 2012 (E-GBM) (1)	None	None	None	None
MS-MLPA	_	_	_	_	_	Park 2011 (2)	None	None	None
PCR-HRM	_	_	_	_		_	Yamashita 2018 (5)	None	None
PCR-mR- NA	_	_	_	_				None	None
Other	_	_	_	_	_	_	_	_	Almuqate 2018 (2)

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(Continued)

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IHC: immunohistochemistry; MS-MLPA: methylation-specific multiplex ligation-dependent probe amplification; MSP: methylation-specific polymerase chain reaction; PCR-HRM: polymerase chain reaction with high-resolution melting; PCR-mRNA: polymerase chain reaction-messenger ribonucleic acid; PSQ: pyrosequencing; qMSP: quantitative methylation-specific polymerase chain reaction.



Appendix 5. Details of methods implemented

Study ID	Method	Technique description	Primers used	Anti- body/mR- NA measure- ment/enzy- matic activity assay	How cut-off threshold deter- mined
Almuqate 2018	Technique: MS- RE-qPCR	MS-RE-qPCR	NR	_	Mention of "opti- mal cut-off;" un-
	Sample type: NR				clear whether the 5% cut-off was prespecified, or
	CpG sites: NR				whether multiple cut-offs were inves-
	Threshold: > 5%				tigated
	Technique: MS- RE-qPCR	-			Described as "cur- rent cut-off" and
	Sample type: NR				"analytically vali- dated"
	CpG sites: NR				
	Threshold: > 9%				
Bady 2012 (E- GBM)	Technique: bead array	Infinium HumanMethylation27 (HM-27K) BeadChip	NR	_	From M-GBM dataset
	Sample type: frozen				
	CpG sites: 31 and 83				
	Threshold: > 0.358				
	Technique: bead array	Infinium HumanMethylation27 beadchip (Illumina Inc.)	NR	_	Selected the threshold that gave
	Sample type: frozen				the best stratifica- tion value accord- ing to the log-rank
	CpG sites: 78– 84				test
	Threshold: > 10%				
	Technique: PSQ	Methylation-specific PSQ per-	_	_	"The percentage
	Sample type: frozen	formed with PyroMark Q96 CpG MGMT kit Qiagen			of MGMT methyla- tion was averaged over the 5 CpG-
	CpG sites: 74– 78				sites interrogated The data was di-

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(Continued)	Threshold: > 7.28%				chotomized into unmethylated and methylated status using an iterative procedure based on segmented re- gression [5]. The optimal cut-off ob- tained was 7.28%, defined as the point where the sum of squares of residuals is minimal."
Bady 2012 (M- GBM)	Technique: bead array Sample type: frozen CpG sites: 31 and 83 Threshold: > 0.358	Infinium HumanMethylation27 (HM-27K) BeadChip	NR	_	Threshold derived empirically to max- imise sensitivity and specificity: "For classification, we used a probabili- ty cut-off of 0.358, which empirically maximized the sum of sensitivity and specificity."
	Technique: MSP Sample type: NR CpG sites: 76– 80 and 84–86 Threshold: NR	"Performed basically as reported by Esteller et al." Esteller M et al. <i>New England Journal of Medicine</i> 2000;343:1350–4.	See Figure 1 of publication	_	NR
Barault 2015	Technique: Methyl-beam- ing Sample type: FFPE CpG sites: 79– 83 Threshold: > 40.2%	Methyl-beaming assay. "BEAM- ing analysis is a multistep digital PCR based technique published by Diehl and colleagues [7]. Its appli- cation for methylation is named Methyl-BEAMing and has been pre- viously described to detect methy- lation of the VIM gene [5]The percentage of methylation was calculated dividing the methy- lated specific signal by the sum of methylated plus unmethylat- ed specific signal." Workflow for methyl-beaming: bisulfite treat- ment; locus enrichment; digital PCR; hybridisation flow cytometry	Methyl-beam- ing 1st PCR: forward 5'-TC- CCGCGAAAT- TAATACGACGTT- TAGGATAT- GTTGGGATAGT-3', reverse 5'-GCTG- GAGCTCTGCAGC- TAAACCACC- CAA-3'. Methyl- beaming emul- sion PCR: for- ward 5'-TCC- CGCGAAATTAAT- ACGAC-3', re- verse 5'-GCTG- GAGCTCTGCAGC- TA-3' (Table S2 of publication).	_	"ROC analysis was carried out to eval- uate the thresh- old best fitting the overall survival (OS) at 1 year" on a co- hort of 98 partic- ipants with GBM diagnosed before TMZ was intro- duced as a compo- nent of standard treatment. The cut- off was then vali- dated in this cohort of participants.
	Technique: PSQ	Bisulfite-PSQ. "Pyrograms were analyzed using PyroMark Q24 Soft- ware, average of the 6 CpG sites	Forward 5'- GTTTAGGATAT- GTTGGGATAGT-3',	_	-

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 97



(Continued)	Sample type: FFPE CpG sites: 76– 81 Threshold: > 29.6%	methylation values was used for further analyses."	reverse 5'- GGACACCGCT- GATCGTTTAAAC- CACCCAAA- CACTCACCAA-3', universal 5'- GGGACACCGCT- GATCGTTTA-3', sequencing 5'-GTTTTTA- GAAYGTTTTGYGTT- T-3' (Table S2 of publication)		
Barbagallo 2014	Technique: MSP Sample type: FFPE CpG sites: 76– 80 and 84–87 Threshold: in- cluding weakly Technique: MSP Sample type: FFPE CpG sites: 76– 80 and 84–87 Threshold: ex- cluding weakly	"MSP assay was performed using a 2-step nested PCR approach as previously described. The MSP reactions were performed in 25 ml by 2720 Thermal Cycler Ap- plied Biosystem PCR. Universal un- methylated and polymethylated DNA were included as controls in each set of reactions, in addition to a negative control sample without DNA."	Primers from Es- teller 1999	_	"Universal un- methylated and polymethylated DNA were includ- ed as controls in each set of reac- tions, in addition to a negative con- trol sample without DNA. Individual tu- mors showing on- ly very weak PCR products for the methylated MGMT sequence promot- er but strong PCR products for the un- methylated MGMT sequence promot- er were judged as "weakly methylat- ed"."
	Technique: PSQ Sample type: FFPE CpG sites: NR Threshold: > 9% Technique: PSQ Sample type: FFPE CpG sites: NR Threshold: > 25%	"Templates for pyrosequencing were amplified with primers that were biotinylated for template strands (MGMT PyroMark CpG As- say kit, Qiagen). The biotinylated polymerase chain reaction (PCR) products were then immobilized on streptavidin-coated Sepharose beads (GE Healthcare), and the sin- gle-stranded DNA templates were analyzed by PyroMark Q24 (Qia- gen)."	Primers from MGMT PyroMark CpG Assay kit, Qiagen		NR
Bell 2017	Technique: QF- IHC (AQUA) Sample type: FFPE	QF-IHC (AQUA). Median cut-off tu- mour mask. 4 tissue microarrays containing paraffin-embedded tumour cores from the 452 RTOG	N/A	Antibody: MGMT (MT3.1) (Santa Cruz; 1:100)	To determine the best cut-off points for markers with continuous values

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(Continued)	CpG sites: N/A	0525 people were cut at 5 μm and			significantly as-
	Threshold: > median	sections were placed on positive- ly charged slides. As a surrogate for tumour colocalisation, proteins were colocalised with glial fibril- lary acidic protein (DAKO;1:100) to stain the cytoplasmic compart- ments of glial cells. Deparaffin- isation and retrieval were per- formed as previously described. Slides were scanned by HistoRx PM-2000 and analysed by AQUA- nalysis software. Each protein was scored in the tumour, cytoplasm, and nuclear components of each tissue microarray core using the HistoRxTM AQUA platform and flu- orescent IHC.			sociated with sur- vival for inclusion in the RPA model, the technique of using ROC curves was ap- plied. Because the area under the ROC curve for all mark- ers was ≤ 0.65, lim- iting the ability to determine optimal cut points, methods using quartiles, ter- tiles and medians were used.
	Technique: qMSP	qMSP assay (detail from the orig- inal NRG RTOG 0525 paper). Per- formed centrally by Oncomethy-	See Vlassen- broeck.	_	From Vlassen- broeck paper: "These cutoffs had
	Sample type: NR	lome Science – direct, real-time MSP (RTOG 0525 Gilbert paper ref-			been defined pre- viously in a small- er data set and are
	CpG sites: NR	erences Vlassenbroeck 2008, MSP method taken from Vlassenbroeck.			consistent with the
	Threshold: > 8	"Validation of Real-Time Methyla- tion-Specific PCR to Determine O6- Methylguanine-DNA Methyltrans- ferase Gene Promoter Methylation in Glioma"). "Analyte (m_MGMT and β -actin [ACTB]) quantification was performed by real-time MSP assays. These consisted of paral- lel amplification/quantification processes using specific primer and primer/detector pairs for each analyte using the Amplifluor as- say format on an ABI Prism 7900HT instrument (Applied Biosystems, Foster City, CA). The analyte de- fined in the direct, real-time MSP was the MGMT promoter sequence and detects the fully methylated version. ACTB was used as a ref- erence gene in the assay, using primers that are outside any CpG islands. The Amplifluor direct for- ward primers are preceded by the detection elements (underlined). The amplicon size is 136 bp for the m_MGMT analyte, including the Amplifluor detection sequence."			present study sug- gesting the cutoff at ratio value 8."
Brigliadori 2016	Technique: PSQ Sample type: FFPE	PSQ. 10 CpG sites of the MGMT promoter (74–83) located in a gene region recognised as critical for transcriptional control (DMR2) were analysed using a PyroMark	Primers that had been biotinylat- ed for template strands (MGMT	_	References litera- ture

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 99 people with glioblastoma treated with temozolomide (Review)

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(Continued)	CpG sites: 74– 83 Threshold: > 9% Technique: PSQ Sample type: FFPE CpG sites: 74– 83 Threshold: > 29%	96 system (Diatech, lesi, Italy), ac- cording to the manufacturer's pro- tocol. All tumour and control sam- ples were analysed in triplicate. Templates for PSQ were ampli- fied using a Rotorgene 6000 with primers that had been biotinylat- ed for template strands (MGMT plus kit, Diatech, lesi, Italy). 20 μL of the biotinylated PCR products were then immobilised on strep- tavidin-coated Sepharose beads (GE Healthcare, Uppsala, Swe- den), and the single-stranded DNA templates were analysed by Py- roMark Q96 system (Diatech, lesi, Italy). Subsequent quantification	plus kit, Diatech, Iesi, Italy).		
		of the methylation density for the 10 investigated CpG sites was per- formed using the PyroMark Q96 software. Methylation percentages for each sample were obtained by calculating the mean of all 10 methylation sites. The median val- ue of the 3 analyses was consid- ered for each methylation level.			
Chai 2018 (7- site cohort)	Technique: PSQ Sample type: frozen CpG sites: 72– 78 Threshold: > 12% Technique: PSQ Sample type: frozen CpG sites: 74– 78 Threshold: > 12% Technique: PSQ Sample type: frozen CpG sites: 75– 78 Threshold: > 12%	"Bisulfite-treated DNA was pre- amplified with the primers (a) F- primer 5'-GTT TYG GAT ATG TTG GGA TAG TT-3'; (b) biotinylated R- primer 5'-biotin-ACR ACC CAA ACA CTC ACC AA-3'. Different samples were analyzed with two indepen- dent assays, and the PSQ primers were (a) 5'-GAT ATG TTG GGA TAG T-3' (for CpGs 72–78) PSQ test- ing was performed on a PyroMark- er Q96 instrument, and the results were analyzed with PyroMarker Q96 software." (Qiagen)	Bisulfite-treat- ed DNA was pre- amplified with the primers (a) F-primer 5'-GTT TYG GAT ATG TTG GGA TAG TT-3'; (b) biotinylat- ed R-primer 5'- biotin-ACR ACC CAA ACA CTC ACC AA-3'. Dif- ferent samples were analysed with 2 indepen- dent assays, and the PSQ primers were (a) 5'-GAT ATG TTG GGA TAG T-3' (for CpGs 72–78).		"We determined the cutoff in this study by similar strategy, compre- hensively consid- ering the ROC like- lihood value, sen- sitivity, specificity, and cutoffs used in reported studies."
Chai 2018 (8- site cohort)	Technique: PSQ	PSQ. CpGs 75–78. "Briefly, bisul- fite-treated DNA was preampli-	Bisulfite-treat- ed DNA was pre-	_	"We determined the cutoff in this

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 100 people with glioblastoma treated with temozolomide (Review)

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(Continued)					
	Sample type: frozen	fied with the primers (a) F-primer 5'-GTT TYG GAT ATG TTG GGA TAG	amplified with the primers (a)		study by similar strategy, compre-
	CpG sites: 75– 78	TT-3'; (b) biotinylated R-primer 5'-biotin-ACR ACC CAA ACA CTC ACC AA-3'. Different samples were	F-primer 5'-GTT TYG GAT ATG TTG GGA TAG TT-3';		hensively consid- ering the ROC like- lihood value, sen-
	Threshold: > 13%	analyzed with two independent assays, and the PSQ primers were(b) 5'-GTT TTT AGA AYG TTT TG-3' (for CpGs 75–82) PSQ test-	(b) biotinylat- ed R-primer 5'- biotin-ACR ACC CAA ACA CTC		sitivity, specificity, and cutoffs used in reported studies."
	Technique: PSQ	ing was performed on a PyroMark- er Q96 instrument, and the results	ACC AA-3'. Dif- ferent samples		
	Sample type: frozen	were analyzed with PyroMarker Q96 software." (Qiagen)	were analysed with 2 indepen- dent assays, and		
	CpG sites: 75– 82		the PSQ primers were(b) 5'-GTT		
	Threshold: > 12%		TTT AGA AYG TTT TG-3' (for CpGs 75–82).		
	Technique: PSQ	-			
	Sample type: frozen				
	CpG sites: 76– 79				
	Threshold: > 11%				
Dahlrot 2018	Technique: DIF	DIF was performed on FFPE tissue on the Autostainer Plus platform (DAKO Denmark A/S, Glostrup, Denmark). Detection was per-	N/A	Antibody: MT23.2; Invit- rogen 1 + 100, CA, USA.	Median value. The AF-all of all MGMT positive nuclei (de- fined as the area of all MGMT-pos- itive nuclei divid-
(NS cohort)	Sample type: FFPE				
	CpG sites: N/A	formed using DAKO CSA II, Bi- otin-Free Catalyzed Amplification			
	Threshold: < 0.2	System (DAKO ref. K1497). Posi- tive controls consisting of tissue cores from different normal and cancer tissues, including 11 high- grade gliomas, were included in each run.			ed by the area of all nuclei), the AF- t of MGMT-positive tumour nuclei (de- fined as the area of MGMT-positive tumour nuclei di- vided by the area of all tumour nu- clei), and the AF-nt of MGMT positive non-tumour nuclei (defined as the area of MGMT positive non-tumour nuclei divided by the area of all non-tumour nuclei) were iden- tified. Only the AF- t of MGMT-positive tumour nuclei (de- fined as the area of MGMT positive tu- mour nuclei divid-

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)



(Continued)

tumour nuclei) was evaluated.

					evaluated.
	Technique: PSQ Sample type: FFPE CpG sites: 74– 78 Threshold: > 9%	PyroMark Q96. MGMT promoter status was determined, measured, established using PSQ (MGMT Py- ro kit; Qiagen, Hilden, Germany). A modified PSQ method published by Collins et al. [29] was used. Af- ter bisulfite conversion of 50–200 ng of DNA using EZ DNA Methyla- tion Kit (Zymo Research, Irvine, CA, USA), nested PCR was carried out with HotStarTaq Master Mix (Qiagen). Obtained PCR product was used as a template in 4 PSQ assays. PSQ was performed on a PyroMark Q96 MD instrument (Qia- gen) using PyroMark Gold Q96 CDT Reagents (Qiagen). as described by the manufacturer.	NR	_	NR
Dahlrot 2018 (RSD cohort)	Technique: DIF Sample type: FFPE CpG sites: N/A Threshold: < 0.2	DIF was performed on formalin fixed paraffin embedded tissue on the Autostainer Plus platform (DAKO Denmark A/S, Glostrup, Denmark). Detection was per- formed using DAKO CSA II, Bi- otin-Free Catalyzed Amplification System (DAKO ref. K1497). Posi- tive controls consisting of tissue cores from different normal and cancer tissues, including 11 high- grade gliomas, were included in each run.	N/A	Antibody: MT23.2; Invit- rogen 1 + 100, CA, USA.	Median value. The AF-all of all MGMT- positive nuclei (de- fined as the area of all MGMT pos- itive nuclei divid- ed by the area of all nuclei), the AF- t of MGMT-positive tumour nuclei (de- fined as the area of MGMT positive tumour nuclei di- vided by the area of all tumour nu- clei), and the AF-nt of MGMT-positive non-tumour nuclei (defined as the area of MGMT-positive non-tumour nuclei divided by the area of all non-tumour nuclei) were iden- tified. Only the AF- t of MGMT-positive tumour nuclei (de- fined as the area of MGMT-positive tu- mour nuclei divid- ed by the area of all tumour nuclei) was evaluated.
	Technique: PSQ Sample type: FFPE	MGMT promoter status was de- termined, measured and estab- lished using PSQ (MGMT Pyro kit; Qiagen, Hilden, Germany) as de-	NR	_	NR

 Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)
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(Continued)	CpG sites: 74–	scribed by the manufacturer. DNA		
	78 Threshold: > 10%	was purified from 10 lm paraffin slides using QIAamp DNA FFPE Tis- sue kit (Qiagen), and MGMT PSQ was performed according to the kit instructions.		
Dunn 2009	Technique: PSQ Sample type: frozen, smear, FFPE or a com- bination CpG sites: 72– 83 Threshold: > 9%	"The pyrosequencing assay was performed as described earlier (Shaw et al, 2006). The primers used for amplification of bisul- phite-treated DNA were forward: 5'-gGGATAGTTGGGATAGTT-3' (the first g avoids formation of hair- pin loops) and reverse: 5'-bi- otin-ATTTGGTGAGTGTTTGGG-3' giving a 99-bp amplicon at ge- nomic position 131 155 467–131 155 565pyrosequencing on a	"The primers — used for am- plification of bisulphite-treat- ed DNA were forward: 5'- gGGATAGTTGGGATAGT- T-3' (the first g avoids formation of hairpin loops) and reverse: 5'- biotin-ATTTG-	≥ mean ± 2 SD for non-neoplastic brain
	Technique: PSQ Sample type: frozen, smear, FFPE, or a com- bination CpG sites: 72– 83 Threshold: > 20%	PSQ96MA System (Biotage, Upp- sala, Sweden) using the primer 5'- GGATATGTTGGGATAGT-3' and Py- roGold reagents (Biotage). The Py- ro Q-CpG software 1.0.9 (Biotage) was used to analyse dataPyrose- quencing yields data for 12 CpG sites within the MGMT promoter. For data analysis, the percentage methylation obtained for each CpG was averaged across the 12 CpGs in duplicate PCR reactions (aver-	GTGAGT- GTTTGGG-3' giv- ing a 99-bp am- plicon at genom- ic position 131 155 467–131 155 565py- rosequencing on a PSQ96MA System (Bio- tage, Uppsala, Sweden) us-	"methylated cas- es were ranked ac- cording to methyla- tion and divided in- to three groups."
	Technique: PSQ Sample type: frozen, smear, FFPE or a com- bination CpG sites: 72– 83 Threshold: > 29%	age methylation per sample)."	ing the primer 5'-GGATAT- GTTGGGATAGT-3'."	"Methylated cases were dichotomised using receiver op- erator character- istic (ROC) plots comparing aver- age methylation per case with the Cox regression sur- vival function for OSReceiver op- erator characteris- tic analysis used to separate methylat- ed cases into two prognostic groups yielded a cut-off of 29.4%."
	Technique: PSQ Sample type: frozen, smear, FFPE or a com- bination CpG sites: 72– 83	-		"methylated cas- es were ranked ac- cording to methyla- tion and divided in- to three groups."

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 Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)
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Threshold: > 35%				
Technique: PSQ				"Unsupervised hi-
Sample type: frozen, smear, FFPE or a com- bination				erarchical cluster analysis of average methylation at each CPG site."
CpG sites: 72– 83				
Threshold: clus- ter 1 vs 2 and 3				
Technique: PSQ				
Sample type: frozen, smear, FFPE or a com- bination				
CpG sites: 72– 83				
Threshold: clus- ter 1 and 2 vs 3				
Technique: IHC	Negative controls were carried out by omission of the primary an- tibodies. Each IHC staining was scored blinded to clinical or mole- cular information. For the assess- ment of MGMT protein expression,	N/A	Antibody:	The fraction of im-
Sample type: FFPE			oclonal anti- body MT 3.1 (Dako).	munopositive tu- mour cells was evaluated semi-
CpG sites: N/A				quantitatively and categorised accord-
Threshold: < 10%	only nuclear staining was consid- ered. Staining of vascular endothe- lial cells served as an internal pos- itive control. The DAKO catalysed signal amplification horseradish peroxidase system was used as the detection systems according to the manufacturer's protocol to show MGMT expression.			ing to the follow- ing immunoreac- tivity scores: 0, no positive tumour cells; 1, weak ex- pression < 10% pos- itive tumour cells; 2, moderate expres- sion 10–50% posi- tive tumour cells; 3, strong expression > 50% positive tu- mour cells.
Technique: MSP Sample type: frozen (14 FF- PE) CpG sites: NR Threshold: NR	Methylation-specific PCR	Methylated MGMT promot- er: 5'-gtttta- gaacgttttgcgtttc- gac-3' and 5'- caccgtccc- gaaaaaaaaactc- cg-3', amplify a 122-bp frag- ment. Unmethy- lated MGMT pro-	_	N/A
	35% Technique: PSQ Sample type: frozen, smear, FFPE or a com- bination CpG sites: 72- 83 Threshold: clus- ter 1 vs 2 and 3 Technique: PSQ Sample type: frozen, smear, FFPE or a com- bination CpG sites: 72- 83 Threshold: clus- ter 1 and 2 vs 3 Technique: IHC Sample type: FFPE CpG sites: N/A Threshold: < 10% Technique: MSP Sample type: frozen (14 FF- PE) CpG sites: NR	35%Technique: PSQSample type: frozen, smear, FFPE or a com- binationCpG sites: 72- 83Threshold: clus- ter 1 vs 2 and 3Technique: PSQSample type: frozen, smear, FFPE or a com- binationCpG sites: 72- 83Threshold: clus- ter 1 and 2 vs 3Technique: IHC Sample type: FFPECpG sites: N/A Threshold: < 10%Threshold: < clust information. For the assess- ment of MGMT protein expression, only nuclear staining was consid- ered. Staining of vascular endothe- lial cells served as an internal pos- itive control. The DAKO catalysed signal amplification horseradish peroxidase system was used as the detection systems according to the manufacturer's protocol to show MGMT expression.Technique: MSPMethylation-specific PCRSample type: frozen (14 FF- PE) CpG sites: NRMethylation-specific PCR	35% Technique: PSQ Sample type: frozen, smear, FFPE or a com- bination CpG sites: 72- 83 Threshold: clus- ter 1 vs 2 and 3 Technique: PSQ Sample type: frozen, smear, FFPE or a com- bination CpG sites: 72- 83 Threshold: clus- ter 1 and 2 vs 3 Technique: IHC Sample type: FFPE Sample type: frozen, smear, FFPE CpG sites: 72- 83 Threshold: clus- ter 1 and 2 vs 3 Technique: IHC Sample type: FFPE CpG sites: N/A Threshold: Image: Sample type: frozen (14 FF- PE) Methylation-specific PCR Methylation-specific PCR MSP Methylation-specific PCR MGMT expression. Technique: MSP Methylation-specific PCR Methylated MGMT peroxidase system was used as the detection systems according to the manufacturer's protocol to show MGMT expression. MSP Sample type: frozen (14 FF- PE) CpG sites: NR Threshold: NR Threshold: NR	35% Technique: PSQ Sample type: frozen, smear, FFPE or a com- bination CpG site: 72- 83 Threshold: clus- ter 1 vs 2 and 3 Technique: PSQ Sample type: frozen, smear, FFPE or a com- bination CpG site: 72- 83 Threshold: clus- ter 1 and 2 vs 3 Technique: IHC Sample type: frozen, smear, FFPE Ngative controls were carried out by omission of the primary an- tibodies. Each IHC staining was scored blinded to clinical or mole- cular information. For the assess- ment of MGMT protein expression, only nuclear staining was consid- ered. Staining of vascular endothe- lial cells served as an internal pos- itive control. The DAKO catalysed signal amplification horseradish peroxidase system was used as the detection systems according to the manufacturer's protocol to show MGMT expression. – MEthylated MGMT promot. er: 5':gtttta- gaaaaaaactc- cg:3', amplify a 122-bp frag- ment. Unmethy-

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)



(Continued)			were 5'-tgtgtttt- tagaatgttttgt- gttttgat-3' and 5'-ctaccaccatcc- caaaaaaaaactc- ca-3', amplify a 129-bp fragment.		
	Technique: PCR-mRNA Sample type: frozen (14 FF- PE) CpG sites: N/A Threshold: < 50%	Expression of MGMT transcripts was determined by real-time re- verse transcription-PCR using the ABI PRISM 5700 sequence detec- tion system (Applied Biosystems). The transcript level of MGMT was normalised to the transcript level of ARF1 (ADP-ribosylation factor 1, GenBank accession-no. M36340).	MGMT-RT-F, 5'-tgcacagc- ctggctgaatg-3' and MGMT-RT- R, 5' ggtgaac- gactcttgctg- gaaa-3' resulting in a 102-bp frag- ment.	mRNA: com- mercially available adult human brain RNA (BD Biosciences) was used as reference for the mRNA ex- pression.	NR
Havik 2012	Technique: MSP Sample type: frozen CpG sites: 76- 80 and 84-87 Threshold: NR	"For MSP, melting curve analysis was used to detect PCR products in our samples (35)Three repli- cates of each sample were used to ensure statistical representativity. Real-time PCR followed by melting curve analysis was run on a CFX96 Touch™ Real- Time PCR Detection system (Bio-Rad Laboratories) Following the last cycle, PCR prod- ucts were incubated for 10 s at 95°C before the melting curve was generated by heating from 65°C to 95°C in increments of 0.5°C/5 s while continuously measuring the fluorescence. The melting curves were analyzed using Bio-Rad CFX Manager Software (Bio-Rad Lab- oratories). Melting peaks deter- mined for methylated and un- methylated controls, respectively, were used to identify methylated and unmethylated PCR products in the samples (EpiTect PCR Control DNA Set, cat. number 59695; Qia- gen). Samples having only methy- lated PCR products and samples having both methylated and un- methylated PCR products were both scored as methylation-posi- tive."	Correspond to those used in Esteller 1999. MSP-MGMT- methylated for- ward 5'-TTTC- GACGTTCGTAG- GTTTTCGC-3', MSP-MGMT- methylated reverse 5'- GCACTCTTC- CGAAAAC- GAAACG-3', MSP-MGMT-un- methylated for- ward 5'-TTTGT- GTTTTGAT- GTTTTGTAG- GTTTTGT-3', MSP-MGMT-un- methylated re- verse 5'-AACTC- CACACTCTTC- CACAACTCTC- CACAAACAAAA- CA-3'.		"Samples having only methylated PCR products and samples having both methylated and unmethylat- ed PCR products were both scored as methylation-posi- tive."
	Technique: PCR-HRM Sample type: frozen CpG sites: 72– 83 Threshold: NR	"Three replicates of each sample were usedReal-time PCR fol- lowed by a melting curve step was run on a CFX96 Touch Real-Time PCR Detection system (Bio-Rad Laboratories)The melting curve step was performed according to the company's recommendation (Bio-Rad Laboratories)The data	MGMT MS- HRM2-forward 5'-GCGTTTCG- GATATGTTGGGA- TA-3', MGMT MS- HRM2-reverse 5'- AACGACCCAAA- CACTCACCAAA-3'	_	NR

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 105 people with glioblastoma treated with temozolomide (Review)



(Continued)

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	files generated by the CFX96 sys- tem were imported using Preci- sion Melt Analysis software (Bio- Rad Laboratories) and further ana- lyzed."			
Technique: PSQ Sample type: frozen CpG sites: 74– 78 Threshold: > 2.68%	PSQ using the PyroMark MD Sys- tem (Qiagen). Bisulfite-treated DNA was amplified in a PCR reac- tion using primers from the Py- roMark Q96 CpG MGMT kit (part number 972032, Qiagen).	NR	_	"The pyrosequenc- ing threshold was determined from the mean methy- lation value in the five analyzed CpG sites and the mean standard devia- tion (X + 2SD) in the four menin- giomas. Glioma samples were scored as methyla- tion positive by py- rosequencing if all five CpG sites had methylation values higher than the re- sulting threshold o 2.68%."
Technique: PSQ Sample type: frozen CpG sites: 74– 78 Threshold: > 6% Technique: PSQ Sample type: frozen CpG sites: 74– 78 Threshold: > 7% Technique: PSQ Sample type: frozen CpG sites: 74– 78	"PyroMark Q96 CpG MGMT kit (cat. number 972032; Qiagen) and the PyroMark MD system (Qiagen)."	PyroMark Q96 CpG MGMT kit (cat. number 972032; Qiagen).		"Receiver operat- ing characteristic (ROC) curve analy- sis was used to es- timate the optimal cut-off value for the two PSQ assays, using the mean percentage MGMT methylation for the CpGs covered by the two assays. The area under the ROG curve (AUROC) was calculated after fit- ting ordinary logis- tic regressions with the dependent var able indicating if a patient lived at least 18 months af- ter diagnosis or no Methylation was in cluded as an inde- pendent variable."

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 106 people with glioblastoma treated with temozolomide (Review)

(Continued)

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1)	Sample type: frozen				
	CpG sites: 74– 78				
	Threshold: > 9%				
	Technique: PSQ	Thera 6%. PSQ. 6% cut-off. "Pyro- Mark therascreen MGMT kit (cat.	PyroMark therascreen	_	"Receiver operat- ing characteristic
	Sample type: frozen	number 971061; Qiagen) and the PyroMark Q24 system (Qiagen)"	MGMT kit (cat- alogue number		(ROC) curve analy- sis was used to es-
	CpG sites: 76– 79		971061; Qiagen)		timate the optimal cut-off value for the two PSQ assays,
	Threshold: > 6%				using the mean percentage MGMT methylation for the
	Technique: PSQ				CpGs covered by the two assays. The
	Sample type: frozen				area under the ROC curve (AUROC) was calculated after fit-
	CpG sites: 76– 79				ting ordinary logis- tic regressions with
	Threshold: > 7%				the dependent vari- able indicating if a patient lived at
	Technique: PSQ				least 18 months af- ter diagnosis or not.
	Sample type: frozen				Methylation was in- cluded as an inde-
	CpG sites: 76– 79				pendent variable."
	Threshold: > 8%				
	Technique: PSQ				
	Sample type: frozen				
	CpG sites: 76– 79				
	Threshold: > 9%				
	Technique: qMSP	"Quantitation of MGMT promoter methylation assessed by qMSP is	MGMT qMSP for- ward primer:	-	Percentage methy- lated reference 0
	Sample type: frozen	described in (34)."	5'-GCGTTTC- GACGTTCG- TAGGT-3', re-		(from Havik 2012) (stated "None" in Johannessen
	CpG sites: 71– 73 and 75–86		verse primer: 5'-CACTCTTC-		2018) "A threshold value for scoring
	Threshold: NR		CGAAAAC- GAAACG-3'. MGMT_1 qMSP forward primer:		methylation posi- tive samples was defined based on the qMSP result of

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)



(Continued)			5'-CGAATAT-		four meningiomas,
			ACTAAAACAAC- CCGCG-3', re- verse primer: 5'- TTTTTTCGGGAGC- GAGGC-3' (as in Havik 2012)		which all had PMR values of zero in both qMSP as- says." (from Havik 2012)
	Technique: qMSP Sample type: frozen CpG sites: 71- 86 Threshold: > 0%	MGMT promoter methylation was quantitatively assessed by 2 qMSP assays, each covering 11 CpG sites (CpGs). The 2 assays analysed CpGs in partially overlapping re- gions (Additional file 1: Figure S1), but detected methylation on op- posite DNA strands. Primers (Med- probe) and 6-FAM labelled minor groove binder (MGB) probes (Ap- plied Biosystems, Life Technolo- gies) were modified from 2 previ- ously reported assays.	qMSP: for- ward primer GCGTTTC- GACGTTCG- TAGGT; re- verse primer CACTCTTCC- GAAAACGAAACG MGMT_1 qMSP: forward primer CGAATATAC- TAAAACAAC- CCGCG; re- verse primer TTTTTCGGGAGC- GAGGC ALU qMSP: forward primer GGT- TAGGTATAGTG- GTTTATATTTG- TAATTTTAG- TA; reverse primer ATTAAC- TAAACTAATCT- TAAACTAATCT- TAAACTCATC- TAAACTCATC- TAAACTCAAC- CTCA.		"Samples with a Ct-value above 35 were censored (re- sulting in a quantity of 0). The percent- age of methylat- ed reference (PMR) was calculated for each sample from the median quan- tity value from the triplicates by di- viding the MGMT/ ALU quantity ra- tio in the target by the MGMT/ALU quantity ratio in the methylated control, and multiplying by 100. A threshold value for scoring methylation posi- tive samples was defined based on the qMSP result of four meningiomas, which all had PMR values of zero in both qMSP assays. Only samples with a PMR value above zero in both as- says were scored as methylation posi- tive."
Hsu 2017 (see Hsu 2015)	Technique: IHC Sample type: FFPE CpG sites: N/A Threshold: < 10%	"Tissue sections were immunos- tained on BOND-MAX immunos- tainer (Leica Microsystems). Nor- mal brain was used as external positive control, a previously proven MGMT methylated GBM was used as negative control."	N/A	Antibody: clone MT3.1 (1:100; Ther- mo, Fremont, CA)	The staining inten- sity of endothe- lial cells was used as a reference for interpretation of positive or nega- tive staining of tu- mour cells. Positive MGMT staining (IHC +) was defined as > 10% of tumour nu- clei with the stain- ing intensity sim- ilar to or slightly weaker than that of the adjacent en-

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 108 people with glioblastoma treated with temozolomide (Review)



(Continued)

				dothelial cells (Fig. 1A). Negative MGMT staining was de- fined as staining that did not fulfil the positive criteria.
Technique: MSP Sample type: FFPE CpG sites: 76– 80 and 84–87 Threshold: NR	1-step MSP was performed as pre- viously described (Hsu 2013).	Unmethylated: USP-F1 5' TTTGT- GTTTTGAT- GTTTTGTAG- GTTTTGT 3' and USP-R1 5' AACTCCA- CACTCTTC- CAAAAACAAAACA 3'; methylated: were MSP-F1 50 TTTCGACGTTCG- TAGGTTTTCGC 30 and MSP-R1 5' GCACTCTTCC- GAAAACGAAACG 3'.		Serial dilutions of the positive control were performed and the lowest concentration of methylated DNA to have a PCR prod- uct was 0.5%. MSP of each sample, in- cluding DNA extrac- tion and bisulfite modification, was performed in dupli- cates in accompa- ny with 100%, 0.5%, and 0% methylat- ed DNA as positive, cut-off, and nega- tive controls in each run. Samples with PCR products of any intensity were regarded as a posi- tive result, whereas those with no PCR products were neg- ative for MSP.
Technique: PSQ Sample type: FFPE CpG sites: 76– 79 Threshold: > 5%	"The methylation status of 4 CpG sites within MGMT promoter re- gion (genomic sequence on chro- mosome 10 from 131,265,519 to 131,265,537: CGACGCC- CGCAGGTCCT CG) was analyzed by therascreen MGMT Pyro Kit (Qia- gen GmbH)."	Primers from the MGMT Pyro Kit.	_	According to the recommendation by the manufactur- er.
Technique: qMSP Sample type: FFPE CpG sites: 77– 80 and 84–87 Threshold: > 0.04%	"The qMSP was performed us- ing QuantiTect SYBR Green PCR Kit (Qiagen GmbH, Hilden, Ger- many) as previously described (Hsu 2015)."	Methylated MGMT-F1 5' TTTCGACGTTCG- TAGGTTTTCGC 3', methylat- ed MGMT-R1 5' GCACTCTTCC- GAAAACGAAACG 3'.	_	Median value based on results of the as- say.
Technique: qMSP				Based on authors previous data.

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)

(Continued)	Sample type: FFPE				
	CpG sites: 77– 80 and 84–87				
	Threshold: > 0.1%				
Karayan- Tapon 2010	Technique: IHC	Percentage of positive cells was determined in the most highly stained areas of each tumour section by counting ≥ 200 contiguous cells. All tumoural cells with nuclear immuno-staining (high or low intensity) were counted as positive.	N/A	Antibody: MT3.1 (Novus	Median value used as cut-off.
12010	Sample type: FFPE			Biologicals)	
	CpG sites: N/A				
	Threshold: < 15.5%				
	Technique: MSP	"The methylation status of the CpG island of MGMT promoter was determined using two-stage PCR [32]."	Study references Palmisano et al.	_	NR
	Sample type: frozen		<i>Cancer Research</i> 60:5954–8 which in turn refer-		
	CpG sites: 76– 80 and 84–87		ences Esteller et al. 1999. Primers for stage 1 (am-		
	Threshold: NR		plification) from Palmisano et al. 2000 MGMT- forward, 5'- GGATATGTTG GGATAGTT-3'; and MGMT- reverse, 5'- CCAAAAACC- CCAAAACC-3'. Primers for stage 2 from Esteller et		
			al. 1999: Primer sequences for the unmethy- lated reaction		
			were 5'-TTTGT- GTTTTGAT- GTTTGTAG- GTTTTTGT-3' (for- ward primer)		
			and 5'-AACTC- CACACTCTTC- CAAAAACAAAA- CA-3' (reverse		
			primer), and for the methylat- ed reaction they were 5'-TTTC- GACGTTCGTAG- GTTTTCGC-3' (for-		
			ward primer) and 5'- GCACTCTTC- CGAAAAC-		

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)



(Continued)

GAAACG-3' (reverse primer).

		verse primer).		
Technique: PCR-mRNA Sample type: frozen CpG sites: N/A Threshold: < 0.39	Quantitative real-time PCR. "RNA (1 μg) was reversed-transcribed using the Superscript II reverse transcriptase (Invitrogen, Carls- bad, CA). The relative expression of MGMT was quantified using the Applied Biosystems TaqMan FAM- labeled probes for MGMT and three housekeeping genes: 18S, RPLPO, and GAPDH. The expression of MGMT in tumors was compared with the expression of MGMT in PBMC (unmethylated DNA) by the 2^-ΔΔCt method [34] using the av- erage Ct of the three housekeeping genes for normalization."	N/A	mRNA: "RNA (1 μ g) was re- versed-tran- scribed us- ing the Su- perscript II reverse tran- scriptase (Invitrogen, Carlsbad, CA). The relative expression of MGMT was quantified us- ing the Ap- plied Biosys- tems Taq- Man FAM-la- beled probes for MGMT and three housekeep- ing genes: 18S, RPLPO, and GAPDH. The expres- sion of MGMT in tumors was com- pared with the expres- sion of MGMT in PBMC (un- methylat- ed DNA) by the 2^- $\Delta\Delta$ Ct method [34] using the av- erage Ct of the three house- keeping genes for normaliza- tion."	Median value used as cut-off.
Technique: PSQ Sample type: frozen	PSQ. CpG 1. "The pyrosequencing methylation assay was performed with the PyroMarkTM MGMT kit (Biotage, Uppsala, Sweden) on a	PyroMarkTM MGMT kit (Bio- tage, Uppsala, Sweden).	_	Median value usec as cut-off.
CpG sites: 74	PSQTM96 MA system (Biotage, Up- psala, Sweden), according to the			
Threshold: > 5.5%	manufacturer's protocol. The Py- roMarkTM MGMT kit detects the level of methylation of five CpG			
Technique: PSQ	sites located in the first exon of the MGMT gene."			
Sample type: frozen	0			

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)

Median value used

as cut-off.

(Continued)

CpG sites: 74– 78

Threshold: > 8.0%

Technique: PSQ

Sample type: frozen

CpG sites: 75

Threshold: > 8.7%

Technique: PSQ

Sample type: frozen

CpG sites: 76

Threshold: > 8.0%

Technique: PSQ

Sample type: frozen

CpG sites: 77

Threshold: > 7.85%

Technique: PSQ

Sample type: frozen

CpG sites: 78

Threshold: > 7.8%

Technique: SQ- MSP	SQ-MSP. "Amplifications were car- ried out on an MX4000 instrument	Study references Palmisano et al.	_
Sample type: frozen	with the Brilliant SYBR Green Core kit (Stratagene, La Jolla, CA) or on an Applied Biosystems ABI-PRISM	<i>Cancer Research</i> 60:5954–8 which in turn refer-	
CpG sites: 76– 80 and 84–87	7900 with the Applera SYBR Green master mix (Applied Biosystems,	ences Esteller et al. 1999. Primers	
Threshold: > 35	Foster City, CA). Methylation index (MI) was calculated using a modi- fication of the formula proposed by Fackler et al.: %M = 100 – [(CtM/ CtM + CtUM) × 100] [33]."	for stage 1 (am- plification) from Palmisano et al. 2000 MGMT- forward, 5'-	
	Ctm + CtOm) ~ 100j [35].	GGATATGTTG GGATAGTT-3'; and MGMT-	

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)

reverse, 5'-CCAAAAACC-



(Continued)			CCAAACCC-3'. Primers for stage 2 from Esteller et al. 1999: Primer sequences for the unmethy- lated reaction were 5'-TTTGT- GTTTTGAT- GTTTTGAT- GTTTTGT-3' (for- ward primer) and 5'-AACTC- CACACTCTTC- CAAAAACAAAA- CA-3' (reverse	
			primer), and for the methylat- ed reaction they were 5'-TTTC- GACGTTCGTAG- GTTTTCGC-3' (for- ward primer) and 5'- GCACTCTTC- CGAAAAC- GAAACG-3' (re- verse primer).	
Kim 2016	Technique: MSP Sample type: FFPE CpG sites: 76- 80 and 84-87 Threshold: NR	SQ-MSP. FFPE, 12% cut-off. "sqMSPCR was performed with primers specific for either "methy- lated" or "unmethylated" DNA. Forward primers were labeled at their 5' end with a fluorescent reporter dye (FAM), as previous- ly described [17]. The PCR prod- ucts corresponding to the "methy- lated" sequences have a size of 82bp while the "unmethylated" sequences have 12 additional nu- cleotides (94bp). Both fragments were amplified in the same reac- tion and PCR products were an- alyzed by capillary electrophore- sis. Estimation of the amount of methylated DNA was calculated with the following formula, ab- breviations are as follows; MF- "methylated" fraction, UM-"un- methylated" fraction: (peak height of the MF/peak height of the UM + MF) × 100." Reference 17: Nguyen et al. <i>Current Cancer Drug Targets</i> 2015;15:624–40.	Study refer- ences Palmisano WA et al. <i>Can- cer Research</i> 60:5954-8 which in turn refer- ences Esteller et al. 1999. Primers for stage 1 (am- plification) from Palmisano et al. 2000 MGMT- forward, 5'- GGATATGTTG GGATATGTTG GGATAGTT-3'; and MGMT- reverse, 5'- CCAAAACC- CCAAACCC-3'. Primers for stage 2 from Esteller et al. 1999: Primer sequences for the unmethy- lated reaction were 5'-TTTGT- GTTTTGAT- GTTTTGAT- GTTTTGAT- GTTTTGAT- GTTTTGT-3' (for- ward primer) and 5'-AACTC- CACCCTCC-	NR

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 113 people with glioblastoma treated with temozolomide (Review)

CACACTCTTC-



(Continued)

Kristensen 2016

		CAAAAACAAAA- CA-3' (reverse primer), and for the methylat- ed reaction they were 5'-TTTC- GACGTTCGTAG- GTTTTCGC-3' (for- ward primer) and 5'- GCACTCTTC- CGAAAAC- GAAACG-3' (re- verse primer).		
Technique: PSQ Sample type: FFPE CpG sites: 74– 78 Threshold: > 9%	PSQ. "The PyroMark Q96 CpG MGMT kit5,10) (Ensembl ID: OT- THUMT00000051009) (Qiagen, Hilden, Germany)PyroGold reagents were used for the PSQ reaction, and the signal was an- alyzed using the PSQ 96MA Sys- tem (Biotage, Uppsala, Sweden). Target CpGs were evaluated by PSQ96MA 2.1 instrument software (Biotage, Uppsala, Sweden)."	PyroMark Q96 CpG MGMT kit (Ensembl ID: OT- THUMT00000051009 (Qiagen, Hilden, Germany).	9)	"Receiver operat- ing characteristic (ROC) curve analy- sis was used to de- termine the cut-off value of mean per- centage of methyla- tion at the five CpGs for predicting the longer survival3). The area under the ROC curve (AUC) was used to deter- mine the optimal threshold of the mean percentage of the methylation at the five CpGs."
Technique: IHC Sample type: FFPE CpG sites: N/A Threshold: at 0%	"Formalin-fixed, paraffin-em- bedded sections were deparaf- finized in xylene and rehydrated in decreasing concentrations of ethanol; Immunoreactivity was vi- sualized with DAB + (DAKO K3468) as chromogen. The immunohis- tochemical reactions were semi- quantitatively evaluated accord- ing to the number of tumor cells stained; For MGMT evaluation, pos- itive endothelial cells, lympho- cytes, and microglia served as pos- itive internal controls."	N/A	Antibody: monoclon- al mouse an- ti-human antibody against MGMT (MAB16200, 1:200, Milli- pore)	NR
Technique: PSQ Sample type: frozen CpG sites: NR Threshold: > 10%	Standard PSQ. "PCR and pyrose- quencing were performed using the Therascreen (R) MGMT Pyro (R) kit according to the manufactur- er's instructions with slight modifi- cations."	Supp Fig 1	_	NR
Technique: qMSP-PSQ	qMSP-PSQ. Quantitative and al- lelic methylation analyses were	Supp Fig 1	_	This technical cut- off was defined fol-

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)



(Continued)	Sample type:	performed using qMSP and melt-			lowing an evalu-
	frozen CpG sites: NR	ing analyses followed by PSQ of methylation-positive sam-			ation of a serial dilution series of
		ples being heterozygous for the			methylated DNA in-
	Threshold: > 0.1%	rs16906252 SNP. Flowchart of method in Fig 1 of the publication.			to unmethylated.
	01270	Sodium bisulfite conversion of the			
		samples was performed using the			
	Technique: qMSP-PSQ	EZ DNA Methylation kit (Zymo Re-			
		search) according to the manu- factures' instructions, with slight			
	Sample type:	modifications; samples were in-			
	frozen	cubated at 42 °C for 30 minutes			
	CpG sites: NR	instead of 37 °C for 15 minutes.			
	Threshold: >	For the bisulfite reaction the alter- native incubation conditions de-			
	5%	scribed in the appendix were used.			
		The LightCycler 480 (Roche Life			
	Technique:	Science) was used for real-time			
	qMSP-PSQ	PCR and melting analysis. The re- al-time PCR cycling protocol start-			
	Sample type:	ed with one cycle of 95 °C for 10			
	frozen	minutes, followed by 40 cycles			
	CpG sites: NR	of 95 °C for 20 seconds, 70 °C for			
	cpo sites. Nit	20 seconds, 72 °C for 20 seconds. The melting step was performed			
	Threshold: >	from 65 °C to 95 °C after a denat-			
	20%	uration step of 1 minute at 95 °C			
		and a hybridization step of 40 °C			
		for 1 minute. For the reaction mix- tures the SYBR Green Master Mix			
		(Roche) was used at a final con-			
		centration at 1×. Final primer con-			
		centrations were 200 nM of each			
		primer, and 25 ng of DNA was used as template. The final reaction vol-			
		ume was 20 µL. Primer sequences			
		have been published previously.			
		The Alu assay used for normalisa-			
		tion was used without the TaqMan probe using an intercalating fluo-			
		rescent dye instead as previously			
		described. PSQ was performed on			
		the PyroMark Q24 (Qiagen) using			
		the PyroMark Gold Q24 reagents (Qiagen), according to the manu-			
		factures' instructions.			
alezari 2013	Technique: IHC	"MGMT immunoreactivity was	N/A	Antibody:	Median value used
		semi-quantitatively assessed by		MT3.1 (Milli-	as cut-off.
	Sample type:	counting the immunostained tu-		pore)	
	FFPE	mor nuclei as a percentage of the			
	CpG sites: N/A	total tumor nuclei…All immuno- histochemical analyses were per-			
	Threshold: <	formed blinded to methylation sta-			
	30%	tus and clinical information."			
	Technique:	MGMT methylation analysis was	First-stage	_	NR
	MSP	performed by MSP according to a	primers (5'-		
		previously published protocol with	GGATAT-		

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 115

(Continued)	Sample type: FFPE CpG sites: 76- 80 and 84-87 Threshold: NR	slight modifications. Samples were subjected to a 2-stage nested PCR strategy using 2 sets of primers.	GTTGGGATAGTT-3' and 5'- CCAAAACCC- CAAACCC-3') and second-stage primers (un- methylated reac- tion: 5'-TTTGT- GTTTG- TA-GGTTTTGT-3' and 5'-AACTC- CACACTCTTC- CAAAAACAAAA- CA-3'; methy- lated reac- tion: 5TTTC- GACGTTCGTAG- GTTTTCGC-3' and 5'- GCACTCTTC- CGAAAAC- GAA-ACG-3').	
	Technique: PSQ Sample type: FFPE CpG sites: 72– 95 Threshold: NR	Bisulfite-modified DNA, generated by the method described above, was used to sequence a portion of the MGMT promoter contigu- ous with and inclusive of the MSP region. Samples were sequenced with a 2-stage nested PCR using the same first-stage primers as those that were used in MSP: 5'- GGATATGTTGGGATAGTT-3' and 5'-CCAAAAACCCCAAACCC-3', and second stage primers 5'-GGATAT- GTTGGGATAGTT-3' and 5'-CACC- TAAAAACACTTAAAAC-3'. The se- quence of each sample was de- termined using Chromas Lite 2.33 (Technelysium Pty Ltd). There did not appear to be any significant difference in yield compared to MSP.	First-stage — primers as those that were used in MSP: 5'-GGATAT- GTTGGGATAGTT-3' and 5'- CCAAAACC- CCAAACCC-3', and second stage primers 5'-GGATAT- GTTGGGATAGTT-3' and 5'-CACC- TAAAAAACACT- TAAAAAC-3'.	Median number of methylated CpG sites used as the threshold defining hypomethylated (< 3 sites) and hy- permethylated (≥ 3 sites)
Lattanzio 2015	Technique: MSP Sample type: FFPE CpG sites: 76– 80 and 84–87 Threshold: NR	MSP using "primers amplifying the exon 1 region of the MGMT gene including the CpG island and sub- sequently the specific primers for either methylated or unmethylat- ed DNA established by Esteller et al (12). Primers used in the first PCR reaction were: 5'-GGATAT- GTTGGGATAGTT-3' (forward primer, GenBank accession num- ber AL355531, nucleotides 46891 to 46908) and 5'-CCAAAACCC- CAAACCC-3' (reverse primer, Gen- Bank accession number AL355531, nucleotides 47162 to 47179) ampli-	"primers am- plifying the ex- on 1 region of the MGMT gene including the CpG island and subsequent- ly the specific primers for ei- ther methylat- ed or unmethy- lated DNA es- tablished by Es- teller et al (12). Primers used	"the results were qualitatively inter- preted as follows: a visible band in the M primer set and absence of the U primer set product indicated a positive methylation status, whereas absence of a M primer set product and pres- ence of a band in the U primer set was evaluated as a

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)



(Continued)

fying a 289-bp fragment...Frozen and FFPE tissue samples were analyzed in triplicate." DNA extracted from snap-frozen samples. in the first PCR reaction were: 5'-GGATAT-GTTGGGATAGT-T-3' (forward primer, Gen-Bank accession number AL355531, nucleotides 46891 to 46908) and 5'-CCAAAAACCC-CAAACCC-3' (reverse primer, GenBank accession number AL355531, nucleotides 47162 to 47179) amplifying a 289bp fragment." Primers from Esteller et al. 1999: primer sequences for the unmethylated reaction were 5'-TTTGT-GTTTTGAT-GTTTGTAG-GTTTTTGT-3' (forward primer) and 5'-AACTC-CACACTCTTC-CAAAAACAAAA-CA-3' (reverse primer), and for the methylated reaction they were 5'-TTTC-GACGTTCGTAG-GTTTTCGC-3' (forward primer) and 5'-GCACTCTTC-CGAAAAC-GAAACG-3' (reverse nrimer)

negative methylation status."

		verse primer).		
Technique:	MSP using "primers amplifying the	"primers am-	_	"the results were
MSP	exon 1 region of the MGMT gene	plifying the ex-		qualitatively inter-
	including the CpG island and sub-	on 1 region of		preted as follows: a
Sample type:	sequently the specific primers for	the MGMT gene		visible band in the
frozen	either methylated or unmethylat-	including the		M primer set and
CpG sites: 76–	ed DNA established by Esteller et	CpG island and		absence of the U
80 and 84–87	al (12). Primers used in the first	subsequent-		primer set product
80 anu 84-87	PCR reaction were: 5'-GGATAT-	ly the specific		indicated a positive
Threshold: NR	GTTGGGATAGTT-3' (forward	primers for ei-		methylation status,
	primer, GenBank accession num-	ther methylat-		whereas absence
	ber AL355531, nucleotides 46891	ed or unmethy-		of a M primer set
	to 46908) and 5'-CCAAAAACCC-	lated DNA es-		product and pres-

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) 117



(Continued)

CAAACCC-3' (reverse primer, Gen-Bank accession number AL355531, nucleotides 47162 to 47179) amplifying a 289-bp fragment...Frozen and FFPE tissue samples were analyzed in triplicate." DNA extracted from snap-frozen samples. tablished by Esteller et al (12). Primers used in the first PCR reaction were: 5'-GGATAT-GTTGGGATAGT-T-3' (forward primer, Gen-Bank accession number AL355531, nucleotides 46891 to 46908) and 5'-CCAAAAACCC-CAAACCC-3' (reverse primer, GenBank accession number AL355531, nucleotides 47162 to 47179) amplifying a 289bp fragment." Primers from Esteller et al. 1999: primer sequences for the unmethylated reaction were 5'-TTTGT-GTTTTGAT-GTTTGTAG-GTTTTTGT-3' (forward primer) and 5'-AACTC-CACACTCTTC-CAAAAACAAAA-CA-3' (reverse primer), and for the methylated reaction they were 5'-TTTC-GACGTTCGTAG-GTTTTCGC-3' (forward primer) and 5'-GCACTCTTC-CGAAAAC-GAAACG-3' (reverse primer).

ence of a band in the U primer set was evaluated as a negative methylation status."

Technique: PSQ	DNA extracted from FFPE samples.	5'-GGATAT-	_	"To determine the
	"PSQ was performed using the Py-	GTTGGGATAGT-		methylation cut-
Sample type:	roMark ID System (Biotage, Upp-	T-3' (forward		off value for PSQ
FFPE	sala, Sweden). The PSQ primers	primer, GenBank		analysis, we ex-
a a 11 - Ta	used for amplification of bisul-	accession num-		tracted DNA from
CpG sites: 72–	fite-treated DNAs were designed	ber AL355531,		a pool of 5 nor-
80	to cover a region including 9 CpG	nucleotides		mal brain tissues
	sites of the MGMT promoter at the	46891 to 46908)		derived from au-
	beginning of the first exon, adja-	and 5'-biotin-		topsies; the aver-

Prognostic value of test(s) for O6-methylguanine–DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)

(Continued)					
	Threshold: ≥ 9%	cent to the region covered by MSP primers (specifically, CpGs 5–6–7– 8–9 in the pyrograms correspond- ed to CpGs included in specific M/ U MSP primers). The primers were 5'-GGATATGTTGGGATAGTT-3' (for- ward primer, GenBank accession number AL355531, nucleotides 46891 to 46908) and 5'-biotin- AC- CCAAACACTCACCAAA-3' (reverse primer, GenBank accession num- ber AL355531, nucleotides 46972 to 46990), which amplified a 99- bp regionPSQ using the forward primer as sequencing primerRe- sulting data were analyzed and quantified with PyroMark CpG Software (Biotage)All samples were analyzed in duplicate."	ACCCAAACACT- CACCAAA-3' (re- verse primer, GenBank acces- sion number AL355531, nu- cleotides 46972 to 46990). For- ward primer also used as the se- quencing primer.		age percentage of methylation of the 5 samples was 8%; thus we considered methylated any tu- mor sample carry- ing ≥9% methyla- tion."
	Technique: PSQ	DNA extracted from snap-frozen			
	Sample type: frozen	samples. PSQ performed as above.			
	CpG sites: 72– 80				
	Threshold: ≥ 9%				
Lechapt-Zal- cman 2012	Technique: IHC Sample type: FFPE	Immunostaining was performed using heat-induced epitope re- trieval, pH 9.0, a labelled method	N/A	Antibody: a mouse prima- ry antibody	This cut-off was the median value of re- activity in the GBM
	CpG sites: N/A	(EnVision Kit; Dako SA, Trappes, France), and automate immunos- tainer (Dako SA) according to the		against MGMT (clone MT3.1; Chemicon	series.
	Threshold: < 15%	manufacturer's protocol. Nega- tive controls consisted of omitting the primary antibody and replac- ing it with an irrelevant antibody of similar isotype. Endothelial stain- ing was used as an internal pos- itive control. A pathologist who was blind to the people' clinical and MGMT methylation data inde- pendently evaluated MGMT stain- ing using a light microscope at 400 magnification. Specimens with- out valid internal positive controls were excluded from the analysis. For each specimen, 5–10 images of representative fields were then ac- quired at 400 magnification. 360– 1790 tumour cells were counted in specimen, and the percentage of positive tumour nuclei was calcu- lated. Endothelial and inflammato- ry cells were excluded from the cell		al, Temecu- la, Calif) was used at 1:200 dilution. mR- NA: NA	

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(Continued)					
	Technique: MSP Sample type: FFPE CpG sites: 76– 80 and 84–87 Threshold: NR	MSP was performed using a 2-step approach. Bisulfite modification of genomic DNA was undertaken by means of the Epitect Kit (Qia- gen SA) according to the manufac- turer's recommendation. PCR am- plification was carried out as de- scribed by Esteller et al. PCR prod- ucts were loaded onto 5% agarose gels, stained with GelRed (Inter- chim, Montlucon, France), and ob- served under ultraviolet illumina- tion.	5'-TTTGT- GTTTTGAT- GTTTTGT-3' (for- ward primer) and 5'-AACTC- CACACTCTTC- CACACTCTTC- CAAAAA CAAAA- CA-3' (reverse primer) for the unmethylat- ed product and 5'-TTTC- GACGTTCGTAG- GTTTTCGC-3' (for- ward primer) and 5'- GCACTCTTCC- GAAAACGAAA CG-3' (reverse primer) for the methylated product.		N/A
McDonald 2013	Technique: MSP Sample type: FFPE CpG sites: 76– 80 Threshold: NR	(From Estellar et al., 1999) 1 μ g of DNA was denatured by sodium hy- droxide and modified by sodium bisulfite. DNA samples were then purified using Wizard DNA purifica- tion resin (Promega), again treat- ed with sodium hydroxide, pre- cipitated with ethanol, and resus- pended in water. Controls without DNA were performed for each set of PCRs. Each PCR reaction (10 μ L) was directly loaded onto nonde- naturing 6% polyacrylamide gels, stained with ethidium bromide, and visualised under ultraviolet il- lumination.	Primer se- quences of MGMT were for the unmethy- lated reac- tion 5'-TTTGT- GTTTTGAT- GTTTTGAT- GTTTTGTAG- GTTTTTGT-3' (up- per primer) and 5'-AACTC- CACACTCTC- CACACTCTC- CACAACACAA		NR
	Technique: PSQ Sample type: FFPE CpG sites: 74– 78 Threshold: > 8%	PSQ. Tumour DNA (500 ng) was bisulphite modified using the EZ DNA methylation kit (Zymo Re- search, Orange CA) according to the manufacturer's recommen- dations. The CpG PSQ methyla- tion assay was performed with the PyroMark MGMT kit (Qiagen) on a PSQe96 MA system (Qiagen) ac- cording to the manufacturer's pro- tocol. Methylation was quantified	NR	_	Determined through a series of segmented re- gressions where the CpG PSQ val- ues were regressed against their rank order. The model with the cut-off of 8% CpG PSQ result- ed in the minimum

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(Continued)		using the Pyromark CpG software (Qiagen).			mean square error and was thus cho- sen (Supplemen- tary Fig 1).
Melguizo 2012	Technique: IHC	Immunostaining was performed using the Bond Polymer Refine	N/A	Antibody: 1:50; Santa	NR
	Sample type: FFPE	Detection system (Leica Microsis- temas S.L.U, Barcelona, Spain).		Cruz Biotech- nology, incm-	
	CpG sites: N/A	Readings were taken automatical- ly with the ACIS III DAKO system for		RNA: NA	
	Threshold: < 25%	quantification IHC and were veri- fied by 2 experienced pathologists.			
	Technique: MSP	Methylation patterns in the CpG island of MGMT were determined	5'-TTTGT- GTTTTGAT-	_	NR
	Sample type: NR	by chemical modification of un- methylated, but not methylated, cytosine to uracil.	GTTTGTAG- GTTTTTGT-3' (for- ward primer) and 5'-AACTC- CACACTCTTC- CACACTCTTC- CAAAAA CAAAA- CA-3' (reverse primer) for the unmethylat- ed product and 5'-TTTC- GACGTTCGTAG- GTTTTCGC-3' (for- ward primer) and 5'- GCACTCTTCC- GAAAACGAAA CG-3' (reverse		
	CpG sites: 76– 80 and 84–87				
	Threshold: NR				
Nguyen 2015	Technique:	FSQ-MS-PCR was using specific	primer) for the methylated product. Unmethylat-	_	Outcome-based
	FSQ-MS-PCR	primers in a semiquantitative mul- tiplexed fluorescent MS-PCR.	ed cytosines, were 5'-TTTGT- GTTTTGAT- GTTTGTAG- GTTTTTGT-3' (for- ward primer, UF) and 5'-AACTC-		approach used. Assessed effect of
	Sample type: frozen or FFPE				multiple cut-off points on survival
	CpG sites: 76– 80 and 84–87				and determined the cut-off point with the best statistical
	Threshold: > 15%		CACACTCTTC- CAAAAAC AAAA- CA-3' (reverse		significance (p val- ue) and the ones associated with
	Technique: FSQ-MS-PCR	_	primer, UR), and the spe- cific primers		the shortest and longest survivals.
	Sample type: frozen or FFPE		for methylat- ed cytosines were 5'-TTTC-		
	CpG sites: 76– 80 and 84–87		GACGTTCGTA GGTTTTCGC-3' (for- ward primer,	-	
	Threshold: > 60%		MF) and 5'- GCACTCTT		

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			CCGAAAAC- GAAACG-3' (re- verse primer, MR).		
Park 2011	Technique: MS- MLPA Sample type: 50% frozen and 50% FFPE CpG sites: NR Threshold: > 0.1% Technique: MS- MLPA Sample type: 50% frozen and 50% FFPE CpG sites: NR Threshold: > 0.2	Used MS-MLPA probe mix pre- pared by MRC-Holland (Salsa MS- MLPA Kit ME011 MMR), which in- cluded 3 probes specific for the MGMT promoter region contain- ing a Hhal recognition site. The procedure was performed accord- ing to the manufacturer's pro- tocol. Hhal (R6441; Promega), a methylation-sensitive restriction enzyme that cuts unmethylated GCGC sites was applied to each set of samples. The resultant PCR frag- ments were separated by capil- lary gel electrophoresis (ABI Prism 7000/7700, Applied Biosystems). The methylation status was quan- tified using GeneMarker software (version 1.5, Soft Genetics). To compensate for the differences in the efficiency of the PCR for the in- dividual samples, the peak value of each probe was normalised by dividing it by the peak of the con- trol probes. To evaluate the methy- lation status, the methylation ra- tio was calculated by the mean of dividing each normalised peak val- ue of the digested sample by that of the corresponding undigested sample. This value corresponds to the percentage of methylated se- quences.	NR		Outcome-based ap- proach: chose the best cut-off to pre- dict early-response evalution (progres- sion/pseudopro- gression).
	Technique: MSP Sample type: 50% frozen and 50% FFPE CpG sites: 76– 80 and 84–86 Threshold: NR	The obtained PCR products were electrophoresed in 2% agarose gels and visualised under ultravio- let illumination after staining with ethidium bromide. For the evalua- tion of the assay results, the prod- ucts from the controls were ex- amined first. The MGMT gene pro- moter fragments in the controls were observed at 80 and 92 bp for the methylated DNA-methylated primer and unmethylated DNA- unmethylated primer combina- tions, respectively. The methylat- ed DNA-unmethylated primer and unmethylated DNA-methylated primer controls were not expected to show any bands. If the control results were acceptable, partici- pant samples were evaluated for the presence of amplification with	The primer se- quences for the MGMT were as follows: methy- lated forward: 5'-TTT CGA CGT TCG TAG GTT TTC GC-3', methy- lated reverse: 5'-GCA CTC TTC CGA AAA CGA AAC G-3', un- methylated for- ward: 5'-TTT GTG TTT TGA TGT TTG TAG GTT TTT GT-3', unmethy- lated reverse: 5'- AAC TCC ACA CTC TTC CAA AAA CAA AAC A-3'.	_	NR

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(Continued)

(Continued)		the methylated and unmethylated primers.			
Quillien 2014 (test)	Technique: IHC Sample type: FFPE CpG sites: N/A Threshold: < 23%	References Chinot et al. <i>Journal of Clinical Oncology</i> 2007;25:1470– 5. Percentage of positive tumour cells determined by a pathologist.	N/A	Antibody: MT-1; Chemi- con, Temecu- la, CA (From Chinot et al. <i>Journal</i> <i>of Clinical</i> <i>Oncology</i> 2007;25:1470– 5, NR in Quil- lien 2012)	"Optimal risk cut- offs were therefore determined as the threshold values of the continuous dis- tribution that best separated low- and high-risk people ac- cording to their out- comes (outcome based method). More precisely, they were defined as the thresholds that op- timized the area un- der the receiver op- erating character- istic (ROC) curve obtained with a Cox model 25 us- ing overall survival (OS) adjusted for age and Karnof- sky score (the pro- portional hazard assumption was checked)."
	Technique: MSP Sample type: frozen CpG sites: 76- 80 and 84-87 Threshold: NR	2-stage PCR	Study references Karayan-Tapon et al. Journal of Neuro-oncolo- gy 2010;97:311– 22, which in turn references Palmisano et al. Cancer Research 20;60:5954–8 which in turn references Es- teller et al. 1999. Primers for stage 1 (ampli- fication) from Palmisano et al. 2000 MGMT- forward, 5'- GGATATGTTG GGATAGTT-3'; and MGMT- reverse, 5'- CCAAAACC- CCAAAACC- CCAAACCC-3'. Primers for stage 2 from Esteller et al. 1999: primer sequences for		NR

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		the unmethy- lated reaction were 5'-TTTGT- GTTTTGAT- GTTTTGTAG- GTTTTTGT-3' (for- ward primer) and 5'-AACTC- CACACTCTTC- CACACTCTTC- CAAAAACAAAA- CA-3' (reverse primer), and for the methylat- ed reaction they were 5'-TTTC- GACGTTCGTAG- GTTTTCGC-3' (for- ward primer) and 5'- GCACTCTTC- CGAAAAC- GAAACG-3' (re- verse primer).	
Technique: Methy- Light-MSP Sample type: frozen CpG sites: 75– 86 Threshold: > 0	MethyLight. Paper cites Metellus P et al. <i>Cancer</i> 2009;115:4783–94. "real-time, fluorescence-based polymerase chain reaction (PCR) was performed using the Light Cy- cler 480 (Roche Diagnostics, Mey- lan, France). Bisulfite-converted genomic DNA was amplified us- ing a set of primers and a fluores- cent dye-labeled oligonucleotide probe, resulting in a semiquanti- tative methylation analysis." In Quillien 2012: "The differences in amounts of input genomic DNA were normalized by the collagen type II, alpha 1 gene (COL2A1). The percentage of methylated refer- ence was calculated as follows: the methylated MGMT/COL2A1 ra- tio for each sample was divided by the same ratio obtained for a SssI- treated genomic DNA used as stan- dard, and values were multiplied by 100."	References Wid- schwendter et al. <i>Cancer Research</i> 2004;64:3807– 13. Forward primer sequence 5'-GCGTTTC- GACGTTCGTAG- GT-3', reverse primer sequence 5'-CACTCTTC- CGAAAAC- GAAACG-3'	"Optimal risk cut- offs were therefore determined as the threshold values of the continuous dis- tribution that best separated low- and high-risk people ac- cording to their out- comes (outcome based method). More precisely, they were defined as the thresholds that op- timized the area un- der the receiver op- erating character- istic (ROC) curve obtained with a Cox model 25 us- ing overall survival (OS) adjusted for age and Karnof- sky score (the pro- portional hazard assumption was checked)."
Technique: PCR-HRM Sample type: frozen CpG sites: 70– 83	"PCR amplification and high-res- olution melting analysis were per- formed using a Mx3000P appara- tus (Stratagene, La Jolla, Calif) (forward: 5' GCGTTTCGGATAT- GTTGGGATAGT 3' and reverse: 5' AACGACCCAAACACTCACCAAA 3') After amplification, a postam-	Forward: 5'- — GCGTTTCG- GATAT- GTTGGGATAGT-3', reverse: 5'-AAC- GACCCAAACACT- CACCAAA-3'.	Melting-curve method. "When the peak corresponding to methylated DNA was >50% of the peak correspond- ing to unmethylat- ed DNA, the sam-

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	Threshold: > 50%	plification melting curve program was initiated by heating to 95°C for 1 minute, cooling to 70°C for 30 seconds, and increasing the temperature to 95°C (heating rate 0.01°C/s) while continuously mea- suring fluorescence. Control DNAs were extracted from blood. The methylated control was obtained by treatment with CpG Methylase M.sssI M0226S (New England Bio- labs, Ipswich, Mass). Sample melt- ing curves were compared with control melting curves obtained with unmethylated and methylat- ed controls. When the peak corre- sponding to methylated DNA was >50% of the peak corresponding to unmethylated DNA, the sample was considered methylated. All re- actions were performed in dupli- cate."		ple was considered methylated."
	Technique: PSQ	"Pyrosequencing was performed with the PyroMark Q96 MGMT kit	PyroMark — Q96 MGMT	"Optimal risk cut- offs were therefore
	Sample type: frozen	(Qiagen, Courtaboeuf, France) on a PSQTM96 MA system (Biotage, Up-	kit (Qiagen, Courtaboeuf,	determined as the threshold values of
	CpG sites: 74	psala, Sweden)."	France) used.	the continuous dis- tribution that best separated low- and high-risk people ac- cording to their out- comes (outcome based method). More precisely, they were defined as the thresholds that op- timized the area un- der the receiver op- erating character-
	Threshold: > 4%			
	Technique: PSQ			
	Sample type: frozen			
	CpG sites: 74– 78			
	Threshold: > 8%			istic (ROC) curve obtained with a
	Technique: PSQ			Cox model 25 us- ing overall survival (OS) adjusted for
	Sample type: frozen			age and Karnof- sky score (the pro-
	CpG sites: 75			portional hazard assumption was
	Threshold: > 11%	_		checked)."
	Technique: PSQ			
	Sample type: frozen			
	CpG sites: 76			

(Continued)	Threshold: > 4%				
	Technique: PSQ				
	Sample type: frozen				
	CpG sites: 77				
	Threshold: > 6%				
	Technique: PSQ				
	Sample type: frozen				
	CpG sites: 78				
	Threshold: > 5%				
	Technique: PSQ	Assay for CpG 74–83. Templates	"forward primer (GTTTYGGATAT- GTTGGGATAG) and a biotiny- lated reverse primer (AAAAC- CACTCRAAAC- TACCAC). Two assays were de- signed and run on this template using two PSQ primers: GAT- AGTTYGYGTTTT- TAGAA (assay for CpGs 74-83) andGYGATTTG- GTGAGTGTTTG (assay for CpGs 84-89)."		"For each of the 16 tested CpG, as well as for the mean of consecutive se- lected CpGs, op- timal risk cut-off was determined as the threshold value of the continuous distribution which best discriminates low- and high-risk people according to their outcomes (outcome-based method). these val- ues were defined as the thresholds that optimized the
	Sample type: frozen	for PSQ were prepared by amplify- ing bisulfite modified DNA with a forward primer (GTTTYGGATATG			
	CpG sites: 74	TTGGGATAG) and a biotinylated re- verse primer (AAAA CCACTCRAAAC-			
	Threshold: > 8%	TACCAC). PSQ was performed us- ing PyroGold Q96 SQA Reagents and the Pyro Q-CpG software on a			
	Technique: PSQ	PyroMark ID pyrosequencer (Qia- gen, Crawley, UK) as per manufac-			
	Sample type: frozen	turer's recommendation. Full de- tails for CpG location and PSQ can be found in Malley et al. [6] and Mullolland et al. [11].			
	CpG sites: 74– 78				
	Threshold: > 9%				
	Technique: PSQ				area under the ROC curve obtained with
	Sample type: frozen				a Cox model [12] using overall sur- vival (OS) and pro-
	CpG sites: 74– 89				gression-free sur- vival (PFS) adjusted for age and Karnof-
	Threshold: > 11%				sky score (the pro- portional hazard assumption was
	Technique: PSQ				checked)."
	Sample type: frozen				
	CpG sites: 75– 79				

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Threshold: > 8%

Technique: PSQ

Sample type: frozen

CpG sites: 76

Threshold: > 5%

Technique: PSQ

Sample type: frozen

CpG sites: 76– 79

Threshold: > 8%

Technique: PSQ

Sample type: frozen

CpG sites: 76– 80

Threshold: > 9%

Technique: PSQ

Sample type: frozen

CpG sites: 77

Threshold: > 7%

Technique: PSQ

Sample type: frozen

CpG sites: 77– 81

Threshold: > 8%

Technique: PSQ

Sample type: frozen

CpG sites: 78

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Threshold: > 4%

Technique: PSQ

Sample type: frozen

CpG sites: 78– 82

Threshold: > 9%

Technique: PSQ

Sample type: frozen

CpG sites: 79

Threshold: > 7%

Technique: PSQ

Sample type: frozen

CpG sites: 79– 83

Threshold: > 8%

Technique: PSQ

Sample type: frozen

CpG sites: 80

Threshold: > 4%

Technique: PSQ

Sample type: frozen

CpG sites: 81

Threshold: > 8%

Technique: PSQ

Sample type: frozen

CpG sites: 82

(Continued)

Threshold: > 16%

Technique: PSQ

Sample type: frozen

CpG sites: 83

Threshold: > 10%

Technique: PSQ

Sample type: frozen

CpG sites: 84

Threshold: > 9%

Technique: PSQ

Sample type: frozen

CpG sites: 84– 88

Threshold: > 17%

Technique: PSQ

Sample type: frozen

CpG sites: 84– 89

Threshold: > 22%

Technique: PSQ

Sample type: frozen

CpG sites: 85

Threshold: > 5%

Technique: PSQ

Sample type: frozen

CpG sites: 85– 89

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(Continued)	Threshold: > 13%			
	Technique: PSQ			
	Sample type: frozen			
	CpG sites: 86			
	Threshold: > 11%			
	Technique: PSQ			
	Sample type: frozen			
	CpG sites: 87			
	Threshold: > 25%			
	Technique: PSQ			
	Sample type: frozen			
	CpG sites: 88			
	Threshold: > 4%			
	Technique: PSQ			
	Sample type: frozen			
	CpG sites: 89			
	Threshold: > 12%			
Quillien 2014	Technique: PSQ	Assay for CpG 74–83. PSQ per-	Forward primer — (GTTTYGGATATG TTGGGATAG) and a biotinylated	— This was the opti-
(validation)	Sample type: FFPE	formed using PyroGold Q96 SQA Reagents and the Pyro Q-CpG soft- ware on a PyroMark ID pyrose-		mal risk cut-off in the initial popula- tion of 89 partici-
	CpG sites: 74– 78	quencer (Qiagen, Crawley, UK) as per manufacturer's recommenda- tion.	reverse primer (AAAA CCACT- CRAAACTACCAC).	pants with GBM.
	Threshold: > 9%			
	Technique: PSQ			
	Sample type: FFPE			
	CpG sites: 74– 78			

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(Continued)	Threshold: > 10%	_			
	Technique: PSQ				
	Sample type: FFPE				
	CpG sites: 74– 78				
	Threshold: > 28%				
Quillien 2016	Technique: PSQ	FFPE, 6% cut-off. "PSQ was per-	PyroMark CpG MGMT kit (ref.	_	Optimised cut- off (current se-
	Sample type: FFPE	formed as previously described [10, 12] using the PyroMark CpG MGMT kit (ref. 972032, Qiagen, France). All assays were performed in duplicate and each result was averaged together. The average percentage of the 5 CpGs tested was considered."	972032, Qiagen, France).		ries/frozen sam- ples). "Optimal risk
	CpG sites: 74– 78				cut-offs were de- termined as pre- viously described
	Threshold: > 6%				with age and per- formance status in- troduced as adjust- ment factors [10]." Reference 10: Quil- lien et al. <i>Cancer</i> 2012;118:4201–11.
	Technique: PSQ				Optimised cut-
	Sample type: FFPE				off (previous se- ries/frozen sam- ples).
	CpG sites: 74– 78				
	Threshold: > 8%				
	Technique: PSQ				Best level of con- cordance between
	Sample type: FFPE				frozen and FFPE samples.
	CpG sites: 74– 78				
	Threshold: > 12%				
	Technique: PSQ				Best level of con-
	Sample type: FFPE				cordance between frozen and FFPE samples.
	CpG sites: 74– 78				
	Threshold: > 13%				

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(Continued)					
	Technique: PSQ Sample type: FFPE				Optimised cut- off (current se- ries/frozen sam- ples). "Optimal risk cut-offs were deter-
	CpG sites: 74– 78				mined as previously described with age
	Threshold: > 16%				and performance status introduced as adjustment fac-
	Technique: PSQ	-			tors [10]." Refer- ence 10: Quillien
	Sample type: frozen				et al. <i>Cancer</i> 2012; 118:4201–11.
	CpG sites: 74– 78				
	Threshold: > 6%				
	Technique: PSQ				Optimised cut- off (previous se-
	Sample type: frozen				ries/frozen sam- ples)
	CpG sites: 74– 78				
	Threshold: > 8%				
	Technique: PSQ	-			Best level of con-
	Sample type: frozen				cordance between frozen and FFPE samples
	CpG sites: 74– 78				
	Threshold: > 12% or 13%				
	Technique: PSQ	PSQ cut-off 12% (Qiagen kit) Hs_MGMT_01_PM PyroMark CpG	PyroMark CpG assay (ref 970032	_	The mean of the methylation at the
	Sample type: frozen	assay (ref 970032 and 972032).	and 972032).		4 CpG sites as pre- defined in previ-
	CpG sites: 74– 78				ous study (Quillien 2014)
	Threshold: > 12%				
	Technique: PSQ	PSQ. Frozen, 16% cut-off. "PSQ	PyroMark CpG	_	Optimised cut-
	Sample type: frozen	was performed as previously de- scribed [10, 12] using the PyroMark CpG MGMT kit (ref. 972032, Qiagen,	MGMT kit (ref. 972032, Qiagen, France).		off (current se- ries/frozen sam- ples). "Optimal risk
	CpG sites: 74-78	France). All assays were performed in duplicate and each result was averaged together. The average			cut-offs were deter- mined as previously described with age and performance

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(Continued)	Threshold: > 16%	percentage of the 5 CpGs tested was considered."		status introduced as adjustment fac- tors [10]." Refer- ence 10: Quillien et al. <i>Cancer</i> 2012; 118:4201–11.
	Technique: PSQ Sample type: frozen CpG sites: 76– 79 Threshold: > 8% Technique: PSQ Sample type: frozen CpG sites: 76– 79 Threshold: > 12%	Therascreen MGMT Pyro Kit (ref. 971061, Qiagen, France) according to the manufacturer's instructions.	N/A —	The mean of the methylation at the 4 CpG sites as pre- defined in previ- ous study (Quillien 2014)
	Technique: SQ- MSP Sample type: FFPE CpG sites: 76– 80 and 84–87 Threshold: > 12%	SQ-MSP. FFPE, 12% cut-off. "sqMSPCR was performed with primers specific for either "methy- lated" or "unmethylated" DNA. Forward primers were labeled at their 5' end with a fluorescent reporter dye (FAM), as previous- ly described [17]. The PCR prod- ucts corresponding to the "methy- lated" sequences have a size of 82bp while the "unmethylated" sequences have 12 additional nu- cleotides (94bp). Both fragments were amplified in the same reac- tion and PCR products were an- alyzed by capillary electrophore- sis. Estimation of the amount of methylated DNA was calculated with the following formula, ab- breviations are as follows; MF- "methylated" fraction: (peak height – of the MF/peak height of the UM + MF) × 100." Reference 17: Nguyen et al. <i>Current Cancer Drug Targets</i> 2015;15:624–40.	"The technique — is using specif- ic primers in a semi-quantita- tive multiplexed fluorescent MS- PCR. Primer se- quences recog- nizing unmethy- lated cytosines were 5'-TTTGT- GTTTTGAT- GTTTTGAG- GTTTTGTAG- GTTTTGTAG- GTTTTGTAG- GTTTTGT-3' (for- ward primer, UF) and 5'-AACTC- CACACTCTTC- CAAAAACAAAA- CA-3' (reverse primer, UR), and the spe- cific primers for methylat- ed cytosines were 5'-TTTC- GACGTTCGTAG- GTTTTCGC-3' (for- ward primer, MF) and 5'- GCACTCTTC- CGAAAAC- GAAACG-3' (re- verse primer,	Best level of con- cordance between frozen and FFPE samples.
	Technique: SQ- MSP Sample type: FFPE CpG sites: 76– 80 and 84–87 Threshold: > 13%			Best level of con- cordance between frozen and FFPE samples.
	Technique: SQ- MSP Sample type: FFPE CpG sites: 76– 80 and 84–87			Optimised cut-off (current series/FF- PE samples). "Op- timal risk cut-offs were determined as previously de- scribed with age and performance status introduced as adjustment

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(Continued)					
	Threshold: > 23% Technique: SQ- MSP Sample type: frozen CpG sites: 76- 80 and 84-87 Threshold: > 13% Technique: SQ- MSP Sample type: frozen CpG sites: 76- 80 and 84-87 Threshold: > 23%	-	MR). The ana- lyzed sequences were designed to target the distal part of the CpG island region of MGMT promot- er, whose com- plete methyla- tion has been correlated with MGMT promoter silencing in can- cer cell lines and primary tumors [18]. Primers were labeled at their 5' end with a fluores- cent reporter dye (FAM)." From: Nguyen et al. <i>Current Cancer</i> <i>Drug Targets</i> 2015;15:624–40.		factors [10]." Ref- erence 10: Quil- lien V et al. <i>Cancer</i> 2012;118:4201–11. Optimised cut- off (current se- ries/frozen sam- ples) and best lev- el of concordance between frozen and FFPE samples. "Optimal risk cut- offs were deter- mined as previous- ly described with age and perfor- mance status in- troduced as adjust- ment factors [10]." Reference 10: Quil- lien et al. <i>Cancer</i> 2012;118:4201–11. Optimised cut-off (current series/FF- PE samples). "Op- timal risk cut-offs were determined as previously de- scribed with age and performance status introduced as adjustment factors [10]." Ref- erence 10: Quil- lien V et al. <i>Cancer</i> 2012;118:4201–11.
Thon 2017	Technique: MSP Sample type: frozen CpG sites: 76– 80 and 84–87 Threshold: NR	Bisulfite conversion of 200–400 ng DNA was performed with the Epi- Tect Bisulfite Kit (Qiagen) as de- scribed previously.	2 pairs of primers, each specific for either the methylated or the unmethy- lated MGMT pro- moter region, were used as originally de- scribed by Es- teller and col- leagues.	_	NR
	Technique: se- quencing Sample type: frozen CpG sites: 75– 99 (unclear)	Taken from Eigenbrod 2014: "The sequencing reaction covers a 316 bp region of the MGMT promoter with 25 CpG sites, including those detected by MSP (corresponding to CpG positions 2–14)."	_	-	"The MGMT pro- moter was consid- ered "methylated" when more than half of the CpG sites (≥13 of the 25 CpG sites) were found to be "methylated"

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(Continued)

Threshold: > 50%

or "partially methylated." A "partially methylated" CpG site was defined as the cytosine peaks being 50 % or more of the corresponding thymine peak. Positions with cytosine peaks as small as 10–50 % of the thymine peak were considered weakly methylated. When 9 – 12 of 25 CpG sites were "methylated/partially methylated" the MGMT promoter was considered "partially" methylated. When more than 9 of the 25 CpG sites were "methylated/partially methylated" the MGMT promoter was considered not methylated." "Sequencing of bisulfite-modified DNA indicated a methylated promoter when more than half of the CpG sites (13 of 25 CpG sites) were found to be methylated; partial methylation was defined as the cytosine and thymine peaks being equally sized or the cytosine peak being twice as high as the corresponding thymine peak."

Yamashita	Technique:	"Converted DNA was subjected	"The primer -	_	Samples with 5%
2018	MSP	to MS-PCR using 2 primer pairs	sequences for		methylation fea-
	_	designed for the amplification of	unmethylat-		tured a faint posi-
	Sample type:	methylated and unmethylated al-	ed reactions		tive band, suggest-
	frozen	leles of the MGMT promoterAm-	was 5'-TTTGT-		ing that the appro-
		plified products were loaded on	GTTTTGAT-		priate cut-off value
	CpG sites: 76–	16% polyacrylamide gels and visu-	GTTTGTAG-		for MS-PCR was 5%.
	80 and 84–87	alized under ultraviolet light using	GTTTTTGT-3' (for-		
	Threshold: NR	ethidium bromide staining."	ward), 5'-AACTC-		
	Theshold. Nix	-	CACACTCTTC-		
			CAAAAACAAAA-		
			CA-3' (reverse);		
			for methylat-		
			ed reactions it		

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frozen

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(Continued)

		was 5'-TTTC- GACGTTCGTAG- GTTTTCGC-3' (for- ward), 5'- GCACTCTTC- CGAAAAC- GAAACG-3' (re- verse)."	
Technique: PCR-HRM Sample type: frozen CpG sites: 72– 89 Threshold: > 5%	StepOne system (Thermo Fisher Scientific). "MS-HRM data were an- alyzed using HRM software (ver- sion 3.0.1, Thermo Fisher Scientif- ic). Output plots were produced as aligned melting curvesThe area under the curve (AUC) was calculated from the aligned melt- ing curves using ImageJ (NIH); lin- ear regression was applied to in-	"The primers — sets were 5'- GCGTTTCG- GATAT- GTTGGGATAGT-3' (for- ward), 5'-CCTA- CAAAACCACTC- GAAACTAC- CA-3' (reverse) primer 1."	Validation of ROC analysis.
Technique: PCR-HRM	terpolate unknown samples from the standardsAll measurements were performed in triplicate."		Validation of ROC analysis.
Sample type: frozen			
CpG sites: 72– 89			
Threshold: > 8%			
Technique: PCR-HRM			From ROC analysis.
Sample type: frozen			
CpG sites: 72– 89			
Threshold: > 10%			
Technique: PCR-HRM			Validation of ROC analysis.
Sample type: frozen			
CpG sites: 72– 89			
Threshold: > 12%			
Technique: PCR-HRM			Validation of ROC analysis.
Sample type: frozen			

 Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)
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(Continued)	CpG sites: 72– 89				
	Threshold: > 15%				
Yang 2012	Technique: IHC	"Staining for MGMT protein on	N/A	Antibody:	NR
	Sample type: FFPE	primary tumor samples was per- formed using anti-MGMT antibody clone MT3.1 (Abcam, Cambridge,		MT3.1 (Ab- cam, Cam- bridge, UK).	
	CpG sites: N/A	England)To calculate the MGMT labeling index of MGMT-positive			
	Threshold: < 10%	cells, the number of immunoreac- tive tumor cells was determined for at least 1,000 cells in randomly selected fields."			
	Technique: MSP	"The converted DNA was subject- ed to methylation-specific PCR us-	"The primer sequences of	_	NR
	Sample type: FFPE	ing two primer sets designed for amplifying the methylated or un- methylated allele of the MGMT	MGMT for the un- methylated and methylated reac-		
	CpG sites: 76– 80 and 84–87	promoterAmplified products were separated on a 3% agarose gel and visualized under UV illumi-	tions were as fol- lows: 5'-TTTGT- GTTTTGAT-		
	Threshold: NR	nation."	GTTTGTAG- GTTTTGT-3' (for- ward) and 5'-AACTCCA- CACTCTTC- CAAAAACAAAA- CA-3' (reverse); and 5'-TTTC- GACGTTCGTAG- GTTTTCGC-3' (for- ward) and 5'- GCACTCTTC- CGAAAAC- GAAACG-3' (re- verse), respec- tively."		
Yoshioka 2018	Technique: SQ- MSP	Brilliant II SYBR Green qPCR Mas- ter Mix and 2 types of primers were used for MSP.	Corresponded to those used in Esteller 1999.	_	Based on Delta Ct values "The ∆Ct val- ues of the tumors
	Sample type: frozen		mMGMT for- ward5'-TTTC-		having no peak at 81° C in dissocia-
	CpG sites: 76– 80 and 84–87		GACGTTCGTAG- GTTTTCGC-3', mMGMTreverse		tion curve were be- tween 4 and 10;" "The smaller the
	Threshold: > 0	_	5'-GCACTCTTC- CGAAAAC-		∆Ct value is, the greater the pro-
	Technique: SQ- MSP		GAAACG-3', uMGMT for- ward 5'-TTTGT-		portion of methy- lated cells and the greater the extent
	Sample type: frozen		GTTTTGAT- GTTTGTAG- GTTTTTGT-3',		of the methylated region in each cell. Therefore, we set
	CpG sites: 76– 80 and 84–87		and uMGMT re- verse 5'-AACTC- CACACTCTTC-		five cutoffs."

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review)

Continued)	Threshold: > 2	CAAAAACAAAA- CA-3'.
	Technique: SQ- MSP	CA-S.
	Sample type: frozen	
	CpG sites: 76– 80 and 84–87	
	Threshold: > 4	
	Technique: SQ- MSP	
	Sample type: frozen	
	CpG sites: 76– 80 and 84–87	
	Threshold: > 6	
	Technique: SQ- MSP	
	Sample type: frozen	
	CpG sites: 76– 80 and 84–87	
	Threshold: > 8	

AF: area fraction; bp: base pair; CpG: 5'-cytosine-phosphate-guanine-3'; DIF: double immunofluorescence; FFPE: formalin-fixed paraffin-embedded; GBM: glioblastoma; IHC: immunohistochemistry; MGMT: O⁶-methylguanine–DNA methyltransferase; mRNA: messenger ribonucleic acid; MS-MLPA: methylation-specific multiplex ligation-dependent probe amplification; MS-RE-qPCR: methylation-specific restriction enzyme quantitative polymerase chain reaction; MSP: methylation-specific polymerase chain reaction; N/A: not applicable; NR: not reported; PCR: polymerase chain reaction; PCR-HRM: polymerase chain reaction with high-resolution melting; PCR-mRNA: polymerase chain reaction-messenger ribonucleic acid; PSQ: pyrosequencing; qMSP: quantitative methylation-specific polymerase chain reaction; qMSP-PSQ: quantitative methylation-specific polymerase chain reaction with pyrosequencing; RNA: ribonucleic acid; ROC: receiver operating characteristic; SD: standard deviation; TMZ: temozolomide.

 Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in
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 people with glioblastoma treated with temozolomide (Review)
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		CpGs analysed (PCR- based tests)	Thresh- old for methy- lated	D4: PF	D4 (justification)	D5: at- trition	D5 (justification)	D6: oth- er PFs	D6 (jus- tifica- tion)	D7: sel. rep.	D7 (justi- fication)
Al- muqate 2018	MS-RE- qPCR	NR	> 5%	Unclear	Cut-off may have been based on perfor- mance.	Unclear	Insufficient informa- tion	Low RoB	No con- cerns	Unclear	Confer- ence ab- stract – li tle infor- mation re ported.
	MS-RE- qPCR	NR	>9%	Low RoB	No concerns	Unclear	Insufficient informa- tion	Low RoB	No con- cerns	Unclear	Confer- ence ab- stract – li tle infor- mation re ported.
Bady 2012 (E- GBM)	Bead ar- ray	31 and 83	> 0.358	Low RoB	No concerns	Low RoB	No missing data	N/A	_	Low RoB	No con- cerns
ODM)	Bead ar- ray	78–84	> 10%	High RoB	Threshold derived from outcome mea- surement.	Low RoB	No missing data	N/A	_	Low RoB	No con- cerns
	PSQ	74–78	> 7.28%	Low RoB	Cut-off does not seem to be determined by performance.	Low RoB	Only 3/50 missing.	N/A	_	Low RoB	No con- cerns
Bady 2012 (M- GBM)	Bead ar- ray	31 and 83	> 0.358	High RoB	Threshold derived from outcome mea- surement.	Low RoB	No missing data	Low RoB	No con- cerns	Low RoB	No con- cerns
	MSP	76–80 and 84– 86	NR	Low RoB	No concerns	Low RoB	No missing data	Low RoB	No con- cerns	Low RoB	No con- cerns
Barault 2015	Methyl- beaming	79–83	> 40.2%	Low RoB	No concerns	Low RoB	No missing data	N/A	_	Unclear	Unclear why ther was no re

Prognostic value of test(s) for OG-methylguanine–DNA methyltransferase (MGM) people with glioblastoma treated with temozolomide (Review) Copyright © 2021 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

Appendix 6. Detailed risk of bias assessments

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Prognostic value of test(s) for O6-methylguanine-DNA methylti	(Continued)											sult for MSP for this cohort of people (MSP in- vestigat- ed in oth- er cohorts of peo- ple in this study).
Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in		PSQ	76-81	> 29.6%	Low RoB	No concerns	Unclear	Missing data for 11/69. Unclear whether there were important differ- ences between those included in the study and those who were not.	N/A		Unclear	Unclear why there is no re- sult for MSP for this cohort of people (MSP in- vestigat- ed in oth- er cohorts of peo- ple in this study).
omoter methy	Barba- gallo 2014	MSP	76–80 and 84– 87	Includ- ing weakly	Low RoB	No concerns	Low RoB	No missing data	Low RoB	No con- cerns	Low RoB	No con- cerns
vlation for pr		MSP	76–80 and 84– 87	Exclud- ing weakly	Low RoB	No concerns	Low RoB	No missing data	Low RoB	No con- cerns	Low RoB	No con- cerns
edicting o		PSQ	NR	> 9%	Low RoB	No concerns	Low RoB	No missing data	Low RoB	No con- cerns	Low RoB	No con- cerns
verall sur		PSQ	NR	> 25%	Low RoB	No concerns	Low RoB	No missing data	Low RoB	No con- cerns	Low RoB	No con- cerns
vival in 140	Bell 2017	QF-IHC (AQUA)	N/A	> medi- an	Unclear	_	Low RoB	No concerns	Low RoB	No con- cerns	Unclear	Multiple HRs are re- ported for different

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(Continued)											overlap- ping sub- groups of this popu- lation.
	qMSP	NR	>8	Unclear	MSP was performed centrally in original RCT.	Unclear	Of the 833 people from the original RCT, 452 had avail- able tissue and un- derwent the microar- ray of 22 proteins. From these, 320 had data for MSP. The study authors did compare OS be- tween the 452 mi- croarray samples and the remainder of the 833 from the RCT cohort that were not included in this secondary analysis – no significant differ- ence. However, there does not appear to be any further exami- nation of missing da- ta.	Low RoB	No con- cerns	Unclear	Multiple HRs are re- ported for different overlap- ping sub- groups of this popu- lation.
Brigli- adori 2016	PSQ	74-83	>9%	Low RoB	No concerns	Low RoB	No concerns	N/A	_	Low RoB	No con- cerns
2010	PSQ	74-83	> 29%	Low RoB	No concerns	Low RoB	No concerns	N/A	_	Low RoB	No con- cerns
Chai 2018 (7- site co- hort)	PSQ	72-78	> 12%	High RoB	Cut-offs appeared to have been selected based on performance (ROC curve analysis, sensitivity, specificity).	Low RoB	people had to have information on methylation status and OS to be includ- ed, and therefore there is no missing data. The bias that selecting on this ba-	N/A	_	Unclear	The whole set of CpGs analysed was cor- related with OS. In addi- tion a sub-

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						sis has already been covered in domain 1.				set that is part of a Qiagen kit was tested, and a third combina- tion. Un- clear why results for other combina- tions not presented.
PSQ	74–78	> 12%	High RoB	Cut-offs appeared to have been selected based on performance (ROC curve analysis, sensitivity, specificity).	Low RoB	People had to have information on methylation status and OS to be includ- ed, and therefore there were no miss- ing data. The bias that selecting on this basis has already been covered in do- main 1.	N/A		Unclear	The whole set of CpGs analysed was cor- related with OS. In addi- tion a sub- set that is part of a Qiagen kit was tested, and a third combina- tion. Un- clear why results for other combina- tions not presented.
PSQ	75–78	> 12%	High RoB	Cut-offs appeared to have been selected based on performance (ROC curve analysis, sensitivity, specificity).	Low RoB	Participants had to have information on methylation status and OS to be includ- ed, and therefore there is no missing	N/A	_	Unclear	The whole set of CpGs analysed was cor- related

(Continued)							data. The bias that selecting on this ba- sis has already been covered in domain 1.				with OS. In addi- tion a sub- set that is part of a Qiagen kit was tested, and a third combina- tion. Un- clear why results for other combina- tions not presented.
Chai 2018 (8- site co- hort)	PSQ	75-78	> 13%	High RoB	Cut-offs appeared to have been selected based on performance (ROC curve analysis, sensitivity, specificity).	Low RoB	People had to have information on methylation status and OS to be includ- ed, and therefore there were no miss- ing data. The bias that selecting on this basis has already been covered in do- main 1.	N/A	_	Unclear	The whole set of CpGs analysed was cor- related with OS. In addi- tion a sub- set that is part of a Qiagen kit was tested, and a third combina- tion. Un- clear why results for other combina- tions not presented.
	PSQ	75–82	> 12%	High RoB	Cut-offs appeared to have been selected based on performance	Low RoB	People had to have information on methylation status	N/A	-	Unclear	The whole set of CpGs

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(Continued)					(ROC curve analysis, sensitivity, specificity).		and OS to be includ- ed, and therefore there were no miss- ing data. The bias that selecting on this basis has already been covered in do- main 1.			analysed was cor- related with OS. In addi- tion a sub- set that is part of a Qiagen kit was tested, and a thirc combina- tion. Un- clear why results for other combina- tions not presented.
	PSQ	76–79	> 11%	High RoB	Cut-offs appeared to have been selected based on performance (ROC curve analysis, sensitivity, specificity).	Low RoB	People had to have information on methylation status and OS to be includ- ed, and therefore there were no miss- ing data. The bias that selecting on this basis has already been covered in do- main 1.	N/A	Unclear	The whole set of CpGs analysed was cor- related with OS. In addi- tion a sub- set that is part of a Qiagen kit was tested, and a thirc combina- tion. Un- clear why results for other combina- tions not presented.

Prognostic value of test(s) for O6-met	(Continued) Dahlrot 2018 (NS cohort)	DIF	N/A	< 0.2	Low RoB	Although no infor- mation was provid- ed about blindness of analysis, all nuclear identification was per- formed automatical- ly so there is reason to assume that the mea- surements ere objec- tives.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
thylguani		PSQ	74–78	> 9%	Low RoB	No concerns	Low RoB	No concerns	N/A	_	Low RoB	No con- cerns
ne-DNA methyltransferase (MGMT)	Dahlrot 2018 (RSD co- hort)	DIF	N/A	< 0.2	Low RoB	Although no infor- mation was provid- ed about blindness of analysis, all nuclear identification was per- formed automatical- ly so there is reason to assume that the mea- surements ere objec- tives.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
) promote		PSQ	74–78	> 10%	Low RoB	No concerns	Low RoB	No concerns	N/A		Low RoB	No con- cerns
Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in	Dunn 2009	PSQ	72–83	> 9%	Low RoB	Cut-off may or may not have been prespec- ified, but it was not data driven (i.e. not based on ROC curve analysis).	Low RoB	Missing data for 6/115 people treated with chemoradiation during the study pe- riod: four had histol- ogy elsewhere and two had inadequate tissue. Median OS for the complete cohort of 115 people was 12.8 months vs 12.4 months in the 109 in- cluded people.	Low RoB	No con- cerns	Low RoB	No con- cerns
lin 145		PSQ	72–83	> 20%	Unclear	Methylated cases were ranked according to methylation and divid-	Low RoB	Missing data for 6/115 people treat- ed with chemoradia-	Low RoB	No con- cerns	Low RoB	No con- cerns

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(Continued)					ed into 3 groups. This does not seem to have been a prespecified analysis, but not data driven.		tion during the study period: 4 had histol- ogy elsewhere and 2 had inadequate tis- sue. Median OS for the complete cohort of 115 people was 12.8 months vs 12.4 months in the 109 in- cluded people.				
	PSQ	72-83	> 29%	High RoB	ROC analysis used to separate cases into 2 prognostic groups.	Low RoB	Missing data for 6/115 people treat- ed with chemoradia- tion during the study period: 4 had histol- ogy elsewhere and 2 had inadequate tis- sue. Median OS for the complete cohort of 115 people was 12.8 months vs 12.4 months in the 109 in- cluded people.	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	72-83	> 35%	Unclear	Methylated cases were ranked according to methylation and divid- ed into 3 groups. This does not seem to have been a prespecified analysis, but not data driven.	Low RoB	Missing data for 6/115 people treat- ed with chemoradia- tion during the study period: 4 had histol- ogy elsewhere and 2 had inadequate tis- sue. Median OS for the complete cohort of 115 people was 12.8 months vs 12.4 months in the 109 in- cluded people.	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	72-83	Cluster 1 vs 2 and 3	Unclear	Methylated cases were ranked according to methylation and divid- ed into 3 groups. This does not seem to have been a prespecified	Low RoB	Missing data for 6/115 people treat- ed with chemoradia- tion during the study period: 4 had histol- ogy elsewhere and 2	Low RoB	No con- cerns	Low RoB	No con- cerns

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(Continued)					analysis, but not data driven.		had inadequate tis- sue. Median OS for the complete cohort of 115 people was 12.8 months vs 12.4 months in the 109 in- cluded people.				
	PSQ	72-83	Cluster 1 and 2 vs 3	Unclear	Methylated cases were ranked according to methylation and divid- ed into 3 groups. This does not seem to have been a prespecified analysis, but not data driven.	Low RoB	Missing data for 6/115 people treat- ed with chemoradia- tion during the study period: 4 had histol- ogy elsewhere and 2 had inadequate tis- sue. Median OS for the complete cohort of 115 people was 12.8 months vs 12.4 months in the 109 in- cluded people.	Low RoB	No con- cerns	Low RoB	No con- cerns
Felsberg 2009	IHC	N/A	< 10%	Low RoB	No concerns	Low RoB	Missing data were due to issues with the IHC staining.	N/A	_	Low RoB	No con- cerns
	MSP	NR	NR	Low RoB	No concerns	Low RoB	No concerns	N/A	_	Low RoB	No con- cerns
	PCR-mR- NA	N/A	< 50%	Unclear	Insufficient informa- tion	Unclear	mRNA data not available for 64% of study cohort. Large amount of missing data but unclear dif- ferences between missing and included participants.	N/A	_	Low RoB	No con- cerns
Havik 2012	MSP	76–80 and 84– 87	NR	Low RoB	No concerns	Low RoB	Missing data vs the population studied in Havik 2012 cov- ered in Domain 1.	N/A	_	Low RoB	No con- cerns
	PCR- HRM	72-83	NR	Unclear	States that there is "no" threshold, al-	Unclear	Missing data for methylation status	N/A	_	Low RoB	No con- cerns

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(Continued)					though there must have been one.		for 11/48 partici-				
	PSQ	74–78	> 2.68%	Low RoB	No concerns	Low RoB	only participants treated with radio- therapy + TMZ in- cluded.	N/A	_	Low RoB	No con- cerns
	PSQ	74-78	> 6%	High RoB	Not prespecified, da- ta driven. "In order to compare the prognos- tic ability of the dif- ferent methods, the optimal cut-off value for PSQ needed to be identified. ROC curve analysis is the method of choice for predict- ing optimal cut-off val- ues (37, 38). The mean percentage methyla- tion of the CpGs ana- lyzed in the two PSQ assays were used in our ROC curve analy- sis, where methyla- tion cut-off scores (1- 15%) were plotted to identify the optimum cut-off value for the prediction of OS of 18 months or more after surgery. The AUROC results, including HR values, are listed in Ta- ble II. The highest val- ues for AUROC were at a cut-off of 7% for PSQ Therascreen and 7 and 8% for PSQ 96."	Low RoB	Missing data vs the population studied in Havik 2012 cov- ered in Domain 1.	N/A		Low RoB	No con- cerns
	PSQ	74–78	> 7%	High RoB	Not prespecified, da- ta driven. "In order to compare the prognos- tic ability of the dif-	Low RoB	Missing data vs the population studied in Havik 2012 cov- ered in Domain 1.	N/A	_	Low RoB	No con- cerns

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(Continued)					ferent methods, the optimal cut-off value for PSQ needed to be identified. ROC curve analysis is the method of choice for predict- ing optimal cut-off val- ues (37, 38). The mean							Library
(Continued)					percentage methyla- tion of the CpGs ana- lyzed in the two PSQ assays were used in our ROC curve analy- sis, where methyla- tion cut-off scores (1– 15%) were plotted to identify the optimum cut-off value for the prediction of OS of 18 months or more after surgery. The AUROC results, including HR values, are listed in Ta- ble II. The highest val- ues for AUROC were at a cut-off of 7% for PSQ Therascreen and 7 and 8% for PSQ 96."							Better health.
	PSQ	74–78	> 8%	High RoB	Not prespecified, da- ta driven. But this was not the optimal cut-off. "In order to compare the prognostic ability of the different meth- ods, the optimal cut- off value for PSQ need- ed to be identified. ROC curve analysis is the method of choice for predicting optimal cut-off values (37, 38). The mean percent- age methylation of the CpGs analyzed in the two PSQ assays were	Low RoB	Missing data vs the population studied in Havik 2012 cov- ered in Domain 1.	N/A	_	Low RoB	No con- cerns	Cochrane Database of Systematic Reviews

(Continued)				used in our ROC curve analysis, where methy- lation cut-off scores (1–15%) were plotted to identify the opti- mum cut-off value for the prediction of OS of 18 months or more after surgery. The AU- ROC results, including HR values, are listed in Table II. The highest values for AUROC were at a cut-off of 7% for PSQ Therascreen and 7 and 8% for PSQ 96."						Cochrane Library Better health.
PSQ	74-78	> 9%	High RoB	Not prespecified, da- ta driven. But this was not the optimal cut-off. "In order to compare the prognostic ability of the different meth- ods, the optimal cut- off value for PSQ need- ed to be identified. ROC curve analysis is the method of choice for predicting optimal cut-off values (37, 38). The mean percent- age methylation of the CpGs analyzed in the two PSQ assays were used in our ROC curve analysis, where methy- lation cut-off scores (1–15%) were plotted to identify the opti- mum cut-off value for the prediction of OS of 18 months or more after surgery. The AU- ROC results, including HR values, are listed in Table II. The highest	Low RoB	Missing data vs the population studied in Havik 2012 cov- ered in Domain 1.	N/A	Low RoB	No con- cerns	Cochrane Database of Systematic Reviews

(Continued)					values for AUROC were at a cut-off of 7% for PSQ Therascreen and 7 and 8% for PSQ 96."						
	PSQ	76-79	> 6%	High RoB	Not prespecified, da- ta driven. But this was not the optimal cut-off. "In order to compare the prognostic ability of the different meth- ods, the optimal cut- off value for PSQ need- ed to be identified. ROC curve analysis is the method of choice for predicting optimal cut-off values (37, 38). The mean percent- age methylation of the CpGs analyzed in the two PSQ assays were used in our ROC curve analysis, where methy- lation cut-off scores (1–15%) were plotted to identify the opti- mum cut-off value for the prediction of OS of 18 months or more after surgery. The AU- ROC results, including HR values, are listed in Table II. The highest values for AUROC were at a cut-off of 7% for PSQ Therascreen and 7 and 8% for PSQ 96."	Low RoB	Missing data vs the population studied in Havik 2012 cov- ered in Domain 1.	N/A		Low RoB	No con- cerns
	PSQ	76–79	>7%	High RoB	Not prespecified, da- ta driven. "In order to compare the prognos- tic ability of the dif- ferent methods, the optimal cut-off value	Low RoB	Missing data vs the population studied in Havik 2012 cov- ered in Domain 1.	N/A	_	Low RoB	No con- cerns

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(Continued)					for PSQ needed to be identified. ROC curve analysis is the method of choice for predict- ing optimal cut-off val- ues (37, 38). The mean percentage methyla- tion of the CpGs ana-						
					lyzed in the two PSQ assays were used in our ROC curve analy- sis, where methyla- tion cut-off scores (1– 15%) were plotted to identify the optimum cut-off value for the prediction of OS of 18 months or more after surgery. The AUROC results, including HR values, are listed in Ta- ble II. The highest val- ues for AUROC were at a cut-off of 7% for PSQ Therascreen and 7 and 8% for PSQ 96."						
	PSQ	76–79	> 8%	High RoB	Not prespecified, da- ta driven. But this was not the optimal cut-off. "In order to compare the prognostic ability of the different meth- ods, the optimal cut- off value for PSQ need- ed to be identified. ROC curve analysis is the method of choice for predicting optimal cut-off values (37, 38). The mean percent- age methylation of the CpGs analyzed in the two PSQ assays were used in our ROC curve analysis, where methy-	Low RoB	Missing data vs the population studied in Havik 2012 cov- ered in Domain 1.	N/A	_	Low RoB	No con- cerns

Prognostic value of test(s) for O6-methylguanin beople with glioblastoma treated with temozol	(Continued)					lation cut-off scores (1–15%) were plotted to identify the opti- mum cut-off value for the prediction of OS of 18 months or more after surgery. The AU- ROC results, including HR values, are listed in Table II. The highest values for AUROC were at a cut-off of 7% for PSQ Therascreen and 7 and 8% for PSQ 96."						Cochrane Library Better health.
Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in 153 people with glioblastoma treated with temozolomide (Review)		PSQ	76–79	> 9%	High RoB	Not prespecified, da- ta driven. But this was not the optimal cut-off. "In order to compare the prognostic ability of the different meth- ods, the optimal cut- off value for PSQ need- ed to be identified. ROC curve analysis is the method of choice for predicting optimal cut-off values (37, 38). The mean percent- age methylation of the CpGs analyzed in the two PSQ assays were used in our ROC curve analysis, where methy- lation cut-off scores (1–15%) were plotted to identify the opti- mum cut-off value for the prediction of OS of 18 months or more after surgery. The AU- ROC results, including HR values, are listed in Table II. The highest values for AUROC were at a cut-off of 7% for	Low RoB	Missing data vs the population studied in Havik 2012 cov- ered in Domain 1.	N/A	Low RoB	No con- cerns	Cochrane Database of Systematic Reviews



(Continued)					PSQ Therascreen and 7 and 8% for PSQ 96."						
	qMSP	71–73 and 75– 86	NR	Low RoB	No concerns	Low RoB	Missing data vs the population studied in Havik 2012 cov- ered in Domain 1.	N/A	_	Low RoB	No con- cerns
	qMSP	71-86	> 0%	Low RoB	No concerns	Low RoB	Only people treated with radiation thera- py + TMZ included.	N/A	_	Low RoB	No con- cerns
Hsu 2017 (see Hsu	IHC	N/A	< 10%	Low RoB	No concerns	Low RoB	No missing data	Low RoB	No con- cerns	Low RoB	No con- cerns
2015)	MSP	76–80 and 84– 87	NR	Low RoB	No concerns	Low RoB	No missing data	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	76–79	> 5%	Low RoB	No concerns	Low RoB	No missing data	Low RoB	No con- cerns	Low RoB	No con- cerns
	qMSP	77–80 and 84– 87	> 0.04%	Low RoB	No concerns	Low RoB	No missing data	Low RoB	No con- cerns	Low RoB	No con- cerns
	qMSP	77–80 and 84– 87	> 0.1%	Low RoB	No concerns	Low RoB	No missing data	Low RoB	No con- cerns	Low RoB	No con- cerns
Karayan- Tapon 2010	IHC	N/A	< 15.5%	Low RoB	No concerns	Low RoB	Missing data for 3/81 people.	N/A	_	Low RoB	No con- cerns
2010	MSP	76–80 and 84– 87	NR	Low RoB	Likely to be based on presence/absence of bands on a gel but threshold not report- ed.	Low RoB	No missing data	N/A	_	Low RoB	No con- cerns
	PCR-mR- NA	N/A	< 0.39	Low RoB	No concerns	Low RoB	Missing data for 1/81 people.	N/A	_	Low RoB	No con- cerns
	PSQ	74	> 5.5%	Low RoB	No concerns	Low RoB	Missing data for 2/81 people.	N/A	_	Low RoB	No con- cerns

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(Continued)	DCO	74 70	. 0.00/		N	L D . D	Missis = data fam 2/01	N1 / A			N
	PSQ	74–78	> 8.0%	Low RoB	No concerns	Low RoB	Missing data for 2/81 people.	N/A	_	Low RoB	No con- cerns
	PSQ	75	> 8.7%	Low RoB	No concerns	Low RoB	Missing data for 2/81 people.	N/A	_	Low RoB	No con- cerns
	PSQ	76	> 8.0%	Low RoB	No concerns	Low RoB	Missing data for 2/81 people.	N/A	_	Low RoB	No con- cerns
	PSQ	77	> 7.85%	Low RoB	No concerns	Low RoB	Missing methylation data for 2/81 people.	N/A	_	Low RoB	No con- cerns
	PSQ	78	> 7.8%	Low RoB	No concerns	Low RoB	Missing data for 2/81 people.	N/A	_	Low RoB	No con- cerns
	SQ-MSP	76–80 and 84– 87	> 35	Low RoB	No concerns	Low RoB	No missing data	N/A	_	Low RoB	No con- cerns
Kim 2016	MSP	76–80 and 84– 87	NR	Low RoB	Likely to be based on presence/absence of bands on a gel but threshold not report- ed.	Low RoB	Data for all people treated with TMZ.	N/A	_	Low RoB	No con- cerns
	PSQ	74–78	> 9%	High RoB	Data driven – based on the results of ROC analysis (although for the whole cohort, not just those treated with TMZ).	Low RoB	Data for all people treated with TMZ.	N/A	_	Low RoB	No con- cerns
Kris- tensen 2016	IHC	N/A	at 0%	Unclear	It is unclear whether the investigators analysing the results were blinded to clini- cal outcomes.	Low RoB	Low proportion with missing data.	N/A	_	Low RoB	No con- cerns
	PSQ	NR	> 10%	Low RoB	No concerns	Unclear	12% missing data.	N/A	_	Low RoB	No con- cerns
	qMSP- PSQ	NR	> 0.1%	Low RoB	No concerns	Low RoB	No missing data	N/A	_	Low RoB	No con- cerns

people with glioblastoma treated with temozolomide (Review) Copyright © 2021 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

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(Continued)	qMSP- PSQ	NR	> 5%	Low RoB	No concerns	Low RoB	No missing data	N/A	_	Low RoB	No con- cerns
	qMSP- PSQ	NR	> 20%	Low RoB	No concerns	Low RoB	No missing data	N/A	_	Low RoB	No con- cerns
Lalezari 2013	IHC	N/A	< 30%	Low RoB	No concerns	Unclear	Missing data on MGMT status for about 15% of the study population. Unclear if there are systematic differ- ences between those with and without da- ta.	Low RoB	No con- cerns	Low RoB	No con- cerns
-	MSP	76–80 and 84– 87	NR	Low RoB	No concerns	Low RoB	MGMT methylation by MSP was mea- sured in 402 peo- ple (missing < 5% of the population). No information about reason for exclusion from this analysis is given.	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	72–95	NR	Low RoB	Cut-off based on medi- an number of methy- lated CpG sites as re- sulted from the analy- sis.	High RoB	MGMT methylation by BiSEQ was mea- sured in 312 peo- ple (> 25% is miss- ing). No informa- tion about reason for exclusion from this analysis is given.	Low RoB	No con- cerns	Low RoB	No con- cerns
Lat- tanzio 2015	MSP	76–80 and 84– 87	NR	Low RoB	No concerns	Unclear	Missing data for 6/46 enrolled people. No information to judge whether there were important differ- ences between par- ticipants who com- pleted the study and those who did not.	N/A	_	Low RoB	No con- cerns

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(Continued)											
	MSP	76–80 and 84– 87	NR	Low RoB	No concerns	Low RoB	Missing data for just 1 participant, and not due to missing data on MGMT sta- tus.	N/A	_	Low RoB	No cor cerns
	PSQ	72–80	≥9%	Low RoB	No concerns	Low RoB	Missing data for just 1 participant, and not due to missing data on MGMT sta- tus.	N/A	_	Low RoB	No cor cerns
	PSQ	72–80	≥9%	Low RoB	No concerns	Low RoB	Missing data for just 1 participant, and not due to missing data on MGMT sta- tus.	N/A	_	Low RoB	No cor cerns
Lechapt- Zalcman 2012	IHC	N/A	< 15%	Low RoB	Single centre for MGMT testing, analysis of tu- mour specimens blind- ed to MSP data and clinical outcomes. Cut- off was defined by the median value of reac- tivity – as done previ- ously by other authors (referenced).	Low RoB	Low proportion of missing data. No in- formation on differ- ences between miss- ing and non-missing people.	N/A		Low RoB	No cor cerns
	MSP	76–80 and 84– 87	NR	Low RoB	Single centre for MGMT testing, analysis of tu- mour specimens blind- ed to participant iden- tity, threshold not re- ported but mentions a "detailed protocol" so I think can assume this included a prespeci- fied cut-off.	Low RoB	Low proportion of missing data in uni- variate analysis. No information on dif- ferences between missing and non- missing people.	N/A	_	Low RoB	No con cerns
McDon-	MSP	76-80	NR	Low RoB	No concerns	Unclear	Insufficient informa-	Low RoB	No con-	Low RoB	No co

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	PSQ	74–78	> 8%	Low RoB	Although cut-off was determined post-hoc, this was only to di- chotomise the data and was not deter- mined based on out- comes, which does not seem like it would in- crease risk of bias.	Low RoB	Only 2/78 missing data.	Low RoB	No con- cerns	Low RoB	No con cerns
Melguizo 2012	IHC	N/A	< 25%	Low RoB	No concerns	Low RoB	Only 2.5% samples missing data.	N/A	_	Low RoB	No con cerns
	MSP	76–80 and 84– 87	NR	Low RoB	Unclear as no informa- tion about threshold.	Low RoB	No concerns	N/A	_	Low RoB	No con cerns
Nguyen 2015	FSQ-MS- PCR	76–80 and 84– 87	> 15%	High RoB	Cut-offs appeared to have been deter- mined based on per- formance.	Low RoB	States that all clinical and molecular data were fully complete in all people.	Low RoB	_	Low RoB	No con cerns
	FSQ-MS- PCR	76–80 and 84– 87	> 60%	High RoB	Cut-offs appeared to have been deter- mined based on per- formance.	Low RoB	States that all clinical and molecular data were fully complete in all people.	N/A	_	Low RoB	No con cerns
Park 2011	MS-ML- PA	NR	> 0.1%	High RoB	Outcome-based cut-off chosen.	Low RoB	No missing data	N/A	_	Low RoB	No con cerns
	MS-ML- PA	NR	> 0.2	High RoB	Outcome-based cut-off chosen.	Low RoB	No missing data	N/A	_	Low RoB	No con cerns
	MSP	76–80 and 84– 86	NR	Low RoB	No concerns	Low RoB	No concerns	N/A	_	Low RoB	No con cerns
Quillien 2014 (test)	IHC	N/A	< 23%	High RoB	Threshold was not prespecified. Cho- sen based on perfor- mance.	Low RoB	1/100 people exclud- ed because of miss- ing results on IHC due to a technical problem during the staining process (this participant was al-	Low RoB	No con- cerns	Low RoB	No con cerns

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						so excluded from all other analyses).				
MSP	76–80 and 84– 87	NR	Low RoB	Likely to be based on presence/absence of bands on a gel but threshold not report- ed.	Low RoB	1/100 people exclud- ed because of miss- ing results on IHC due to a technical problem during the staining process (this participant was al- so excluded from all other analyses).	Low RoB	No con- cerns	Low RoB	No con cerns
Methy- Light-MSF	75–86	> 0	High RoB	Threshold was not prespecified. Cho- sen based on perfor- mance.	Low RoB	1/100 people exclud- ed because of miss- ing results on IHC due to a technical problem during the staining process (this participant was al- so excluded from all other analyses).	Low RoB	No con- cerns	Low RoB	No con cerns
PCR- HRM	70-83	> 50%	Low RoB	No concerns	Low RoB	1/100 people exclud- ed because of miss- ing results on IHC due to a technical problem during the staining process (this participant was al- so excluded from all other analyses).	Low RoB	No con- cerns	Low RoB	No con cerns
PSQ	74	> 4%	High RoB	Threshold was not prespecified. Cho- sen based on perfor- mance.	Low RoB	1/100 people exclud- ed because of miss- ing results on IHC due to a technical problem during the staining process (this participant was al- so excluded from all other analyses).	Low RoB	No con- cerns	Low RoB	No con cerns

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P	vSQ	74	>8%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con cerns
Ρ	νSQ	74–78	> 8%	High RoB	Threshold was not prespecified. Cho- sen based on perfor- mance.	Low RoB	1/100 people exclud- ed because of miss- ing results on IHC due to a technical problem during the staining process (this participant was al- so excluded from all other analyses).	Low RoB	No con- cerns	Low RoB	No con cerns
P	vsQ	74–78	> 9%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No cor cerns
P	vSQ	74-89	>11%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No cor cerns
P	νSQ	75	> 11%	High RoB	Threshold was not prespecified. Cho- sen based on perfor- mance.	Low RoB	1/100 people exclud- ed because of miss- ing results on IHC due to a technical problem during the staining process (this participant was al- so excluded from all other analyses).	Low RoB	No con- cerns	Low RoB	No cor cerns
P	vSQ	75–79	> 8%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No cor cerns
P	νSQ	76	> 4%	High RoB	Threshold was not prespecified. Cho- sen based on perfor- mance.	Low RoB	1/100 people exclud- ed because of miss- ing results on IHC due to a technical problem during the staining process (this participant was al- so excluded from all other analyses).	Low RoB	No con- cerns	Low RoB	No cor cerns
P	vsQ	76	> 5%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No cor cerns

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	PSQ	76–79	> 8%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
·	PSQ	76–80	> 9%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	77	> 6%	High RoB	Threshold was not prespecified. Cho- sen based on perfor- mance.	Low RoB	1/100 people exclud- ed because of miss- ing results on IHC due to a technical problem during the staining process (this participant was al- so excluded from all other analyses).	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	77	> 7%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	77-81	> 8%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	78	> 4%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	78	> 5%	High RoB	Threshold was not prespecified. Cho- sen based on perfor- mance.	Low RoB	1/100 people exclud- ed because of miss- ing results on IHC due to a technical problem during the staining process (this participant was al- so excluded from all other analyses).	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	78-82	> 9%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	79	> 7%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	79–83	>8%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns

Prognostic value of test(s) for O6-methylguanine-DNA methyltransferase (MGMT) promoter methylation for predicting overall survival in people with glioblastoma treated with temozolomide (Review) Copyright © 2021 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

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	PSQ	80	> 4%	High	Cut-off of methylation	Low RoB	No concerns	Low RoB	No con-	Low RoB	No con-
				RoB	based on outcome.				cerns		cerns
	PSQ	81	> 8%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	82	> 16%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	83	> 10%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	84	> 9%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	84–88	> 17%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	84–89	> 22%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	85	> 5%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	85–89	> 13%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	86	> 11%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	87	> 25%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	88	> 4%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	89	> 12%	High RoB	Cut-off of methylation based on outcome.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
illien L4 lida-	PSQ	74–78	> 9%	Low RoB	Cut-off of methylation based on outcome of testing cohort.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns

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(Continued)	PSQ	74–78	> 10%	Low RoB	Cut-off of methylation based on outcome of testing cohort.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	74–78	> 28%	Low RoB	Cut-off of methylation based on outcome of testing cohort.	Low RoB	No concerns	Low RoB	No con- cerns	Low RoB	No con- cerns
Quillien 2016	PSQ	74–78	> 6%	High RoB	Data driven – based on the results of ROC analysis.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
-	PSQ	74–78	>8%	Low RoB	Threshold found in a previous publication.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	74–78	> 12%	Unclear	Unclear why this was chosen/no justifica- tion.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	74–78	> 13%	Unclear	Unclear why this was chosen/no justifica- tion.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	74–78	> 16%	High RoB	Data driven – based on the results of ROC analysis.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	74–78	> 6%	High RoB	Data driven – based on the results of ROC analysis.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns

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	PSQ	74–78	> 8%	Low RoB	Threshold found in a previous publication.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	74–78	> 12% or 13%	Unclear	Unclear why this was chosen/no justifica- tion.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	74–78	> 12%	Low RoB	From Quillien 2016: Standard Operating Procedure and 10 quality control sam- ples for the determi- nation of MGMT pro- moter methylation were sent to the dif- ferent centres as way of standardisation of the process through- out the multiple cen- tres. This approach re- duced risk of bias due to different setting for prognostic factor mea- surement. Cut-off of methylation based on outcome.	Unclear	From the original co- hort, 10 people who had successful ini- tial PSQ could not have the Thera PSQ. Therefore, the num- ber of people includ- ed in this analysis was 102, but should have been 112. Au- thors commented in paper: "These da- ta are almost identi- cal to those obtained for the overall pop- ulation (n=112 peo- ple, 49%, 44% and AUCROC values of 0.69 and 0.70), indi- cating the absence of bias in the selection of the 102 people for the present cohort."	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	74–78	> 16%	High RoB	Data driven – based on the results of ROC analysis.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
	PSQ	76–79	> 8%	Low RoB	Analysis of intralabo- ratory reproducibility of Thera showed high-	Unclear	From the original co- hort, 10 people who had successful ini-	Low RoB	No con- cerns	Low RoB	No con- cerns

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(Continued)					ly reproducibility of re- sults from the different centres.		tial PSQ could not have the Thera PSQ. Therefore, the num- ber of people includ- ed in this analysis was 102, but should have been 112. Au- thors commented in paper: "These da- ta are almost identi- cal to those obtained for the overall pop- ulation (n=112 peo- ple, 49%, 44% and AUCROC values of 0.69 and 0.70), indi- cating the absence of bias in the selection of the 102 people for the present cohort."				
	PSQ	76–79	> 12%	Low RoB	Analysis of intralabo- ratory reproducibility of Thera showed high- ly reproducibility of re- sults from the different centres.	Unclear	From the original co- hort, 10 people who had successful ini- tial PSQ could not have the Thera PSQ. Therefore, the num- ber of people includ- ed in this analysis was 102, but should have been 112. Au- thors commented in paper: "These da- ta are almost identi- cal to those obtained for the overall pop- ulation (n=112 peo- ple, 49%, 44% and AUCROC values of 0.69 and 0.70), indi- cating the absence of bias in the selection of the 102 people for	Low RoB	No con- cerns	Low RoB	No con- cerns

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(Continued)											
	SQ-MSP	76–80 and 84– 87	> 12%	Unclear	Unclear why this was chosen/no justifica- tion.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
	SQ-MSP	76–80 and 84– 87	>13%	Unclear	Unclear why this was chosen/no justifica- tion.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
	SQ-MSP	76–80 and 84– 87	>23%	High RoB	Data driven – based on the results of ROC analysis.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
	SQ-MSP	76–80 and 84– 87	> 13%	High RoB	Data driven – based on the results of ROC analysis. Although also the cut-off that corre- sponds to best concor- dance which we rated as unclear elsewhere.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
	SQ-MSP	76–80 and 84– 87	> 23%	High RoB	Data driven – based on the results of ROC analysis.	Unclear	Numerous missing data but unclear if those with miss- ing data different to those without.	Low RoB	No con- cerns	Low RoB	No con- cerns
Thon 2017	MSP	76–80 and 84– 87	NR	Low RoB	MSP is usually eval- uated by visibility of a band indicating methylation, there- fore, we judged as pre- specified threshold in the absence of a clear description in the text.	Low RoB	1 participant lost to follow-up after 6 months.	Low RoB	No con- cerns	Low RoB	No con- cerns
	Se- quenc- ing	75–99 (unclear)	> 50%	Low RoB	No concerns	Low RoB	1 participant lost to follow-up after 6 months.	Low RoB	No con- cerns	Low RoB	No con- cerns

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(Continued) Ya- mashita 2018	MSP	76–80 and 84– 87	NR	Low RoB	No cut-off/presence or absence of band.	Low RoB	1 participant lost to follow-up after 6 months.	High RoB	Model includ- ed oth- er MGMT status using al- ternative method.	Low RoB	No con- cerns
	PCR- HRM	72–89	> 5%	High RoB	Data driven – based on the results of ROC analysis. This was not the optimal cut-off.	Low RoB	No missing data for MGMT status or OS. Multivariate analy- ses reportedly for all 75 participants, al- though 1 participant with missing IDH1 status.	N/A	_	High RoB	2 other primer sets were used for PCR-HRM. Only have ROC curve data for these (dis- carded as not as pre- dictive as primer set 1).
	PCR- HRM	72–89	> 8%	High RoB	Data driven – based on the results of ROC analysis. This was not the optimal cut-off.	Low RoB	No missing data for MGMT status or OS. Multivariate analy- ses reportedly for all 75 participants, al- though 1 participant with missing IDH1 status.	N/A	_	High RoB	2 other primer sets were used for PCR-HRM. Only have ROC curve data for these (dis- carded as not as pre- dictive as primer set 1).
	PCR- HRM	72–89	>10%	High RoB	Data driven – based on the results of ROC analysis.	Low RoB	No missing data for MGMT status or OS. Multivariate analy- ses reportedly for all 75 participants, al- though 1 participant	High RoB	Model includes other MGMT status using al-	High RoB	2 sets were used for PCR- HRM. Only have ROC curve data

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(Continued)							with missing IDH1 status.		ternative method		for these (discarded as not as predictive as primer set 1).
	PCR- HRM	72–89	> 12%	High RoB	Data driven – based on the results of ROC analysis. This was not the optimal cut-off.	Low RoB	No missing data for MGMT status or OS. Multivariate analy- ses reportedly for all 75 participants, al- though 1 participant with missing IDH1 status.	N/A		High RoB	2 other primer sets were used for PCR-HRM. Only have ROC curve data for these (dis- carded as not as pre- dictive as primer set 1).
	PCR- HRM	72–89	> 15%	High RoB	Data driven – based on the results of ROC analysis. This was not the optimal cut-off.	Low RoB	No missing data for MGMT status or OS. Multivariate analy- ses reportedly for all 75 participants, al- though 1 participant with missing IDH1 status.	N/A		High RoB	2 other primer sets were used for PCR-HRM. Only have ROC curve data for these (dis- carded as not as pre- dictive as primer set 1).
Yang 2012	IHC	N/A	< 10%	Low RoB	To a degree we can set the cut-off in this study.	Low RoB	No missing data	Low RoB	No con- cerns	Unclear	HRM analyses also per- formed but ex- tractable data not presented.



(Continued)											
	MSP	76–80 and 84– 87	NR	Low RoB	To a degree we can set the cut-off in this study.	Low RoB	No missing data	Low RoB	No con- cerns	Unclear	HRM analyses also per- formed but ex- tractable data not presented.
Yoshioka 2018	SQ-MSP	76–80 and 84– 87	> 0	Low RoB	No concerns	Unclear	No information re- garding 4 missing samples.	Low RoB	No con- cerns	Low RoB	No con- cerns
	SQ-MSP	76–80 and 84– 87	>2	Low RoB	No concerns	Unclear	No information re- garding 4 missing samples.	Low RoB	No con- cerns	Low RoB	No con- cerns
	SQ-MSP	76–80 and 84– 87	> 4	Low RoB	No concerns	Unclear	No information re- garding 4 missing samples.	Low RoB	No con- cerns	Low RoB	No con- cerns
	SQ-MSP	76–80 and 84– 87	> 6	Low RoB	No concerns	Unclear	No information re- garding 4 missing samples.	Low RoB	No con- cerns	Low RoB	No con- cerns
	SQ-MSP	76–80 and 84– 87	> 8	Low RoB	No concerns	Unclear	No information re- garding 4 missing samples.	Low RoB	No con- cerns	Low RoB	No con- cerns

AUROC: area under receiver operating characteristic; CpG: 5'-cytosine-phosphate-guanine-3'; DIF: double immunofluorescence; FSQ-MS-PCR: fluorescent semi-quantitative methylation-specific polymerase chain reaction; GBM: glioblastoma; HR: hazard ratio; IDH: isocitrate dehydrogenase; IHC: immunohistochemistry; MGMT: O⁶-methylguanine–DNA methyltransferase; mRNA: messenger ribonucleic acid; MS-MLPA: methylation-specific multiplex ligation-dependent probe amplification; MS-RE-qPCR: methylation-specific restriction enzyme quantitative polymerase chain reaction; MSP: methylation-specific polymerase chain reaction; N/A: not applicable; NR: not reported; OS: overall survival; PCR: polymerase chain reaction; PCR-HRM: polymerase chain reaction with high-resolution melting; PCR-mRNA: polymerase chain reaction-messenger ribonucleic acid; PF: prognostic factor; PSQ: pyrosequencing; QF-IHC: quantitative fluorescence immunohistochemistry; qMSP: quantitative methylation-specific polymerase chain reaction; qMSP-PSQ: quantitative methylation-specific polymerase chain reaction with pyrosequencing; RCT: randomised controlled trial; RoB: risk of bias; ROC: receiver operating characteristic; sel. rep.: selective reporting; SQ-MSP: semi-quantitative methylation-specific polymerase chain reaction; TMZ: temozolomide.

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Study ID	Domain 1: partic- ipant se- lection	Domain 1 (justi- fication)	Domain 2: sub- sequent treatment	Domain 2 (jus- tification)	Domain 3: outcome measure- ment	Domain 3 (justifi- cation)	Domain 4: prognos- tic factor measure- ment	Domain 4 (justifica- tion)	Domain 6: adjust- ment for other prognos- tic factors	Domain 6 (justifica- tion)
Almuqate 2018	Unclear concerns	Conference ab- stract: little infor- mation	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns
Bady 2012 (E-GBM)	Low con- cerns	No concerns	Low con- cerns	People received Stupp	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Bady 2012 (M-GBM)	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Barault 2015	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Barbagallo 2014	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns
Bell 2017	Low con- cerns	No concerns	Low con- cerns	No concerns	Unclear concerns	Note that OS was measured from a later start point than other papers	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns
Brigliadori 2016	Unclear concerns	People undergo- ing biopsy were not included in our analysis due to the way data were presented (they are not in- cluded in the KM- curve)	Low con- cerns	People received Stupp	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Chai 2018 (7-site co- hort)	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ.	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns

Appendix 7. Applicability assessments

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Chai 2018 (8-site co- hort)	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Dahlrot 2018 (NS cohort)	Low con- cerns	No concerns	Low con- cerns	No concerns	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns
Dahlrot 2018 (RSD cohort)	Low con- cerns	No concerns	Low con- cerns	Not all people received TMZ but only people that received TMZ are includ- ed in the analy- sis	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Dunn 2009	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns
Felsberg 2009	High con- cerns	Study cohort included on- ly people who were treated with open resec- tion and who re- ceived ≥ 2 cycles of chemotherapy, thus excluding people with tu- mour biopsy only or too poor clini- cal condition for chemotherapy, or both	Low con- cerns	No concerns	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Havik 2012	Low con- cerns	No concerns	Low con- cerns	Only people treated with ra- diotherapy + TMZ were in- cluded in the analysis	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns

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Hsu 2017 (see Hsu 2015)	Low con- cerns	No concerns	Low con- cerns	TMZ with con- comitant radio- therapy for all people	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns
Karayan- Tapon 2010	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Kim 2016	Low con- cerns	No concerns	Low con- cerns	No concerns	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Kristensen 2016	Low con- cerns	All people have glioblastoma	Low con- cerns	People received Stupp	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Lalezari 2013	Low con- cerns	1/418 partici- pants did not re- ceive TMZ; 2 died before TMZ	Low con- cerns	Nearly all peo- ple received TMZ, only 1 sur- viving partici- pant did not	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns
Lattanzio 2015	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Lechapt- Zalcman 2012	Low con- cerns	People had TMZ but additionally got Gliadel	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
McDonald 2013	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ.	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Melguizo 2012	Unclear concerns	People had to have KPS > 60 to be included	Low con- cerns	All people re- ceived concur- rent TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Nguyen 2015	Low con- cerns	No concerns	Low con- cerns	No concerns	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns
Park 2011	Low con- cerns	No concerns	Low con- cerns	No concerns	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Quillien 2014 (test)	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns

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Quillien 2014 (vali- dation)	Low con- cerns	No concerns	Low con- cerns	No concerns	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns
Quillien 2016	Low con- cerns	Frozen samples with a histologi- cally estimated tumour cell con- tent < 40% were excluded from the study	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns
Thon 2017	Unclear concerns	Special popula- tion: unsuitable for GTR	Low con- cerns	All people were assigned to re- ceive radiother- apy + TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns
Yamashita 2018	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Unclear concerns	Unclear whether putting multiple methods of deter- mining MGMT sta tus into the mode together would thi reflect rea practice
Yang 2012	Low con- cerns	No concerns	Low con- cerns	All people re- ceived TMZ	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	N/A	No con- cerns
Yoshioka 2018	Low con- cerns	No concerns	Low con- cerns	No concerns	Low con- cerns	Outcome was all- cause mortality	Low con- cerns	No con- cerns	Low con- cerns	No con- cerns

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Appendix 8. Hazard ratios for all methods examined in included studies

Study	Technique	Sample type	CpGs analysed (PCR- based tests)	Threshold for methy- lated	HR (95% confidence interval)	Data source
Almuqate 2018	MS-RE- qPCR	NR	NR	> 5%	4.17 (1.78 to 9.75)	Adjusted HR (for age, sex and surgery)
	MS-RE- qPCR	NR	NR	> 9%	3.57 (1.67 to 7.62)	Adjusted HR (for age, sex and surgery)
Bady 2012 (E-GBM)	Bead array	Frozen	31 and 83	> 0.358	3.28 (1.68 to 6.41)	Unadjusted HR
	Bead array	Frozen	78-84	> 10%	5.56 (1.25 to 25.0)	Unadjusted HR
	PSQ	Frozen	74–78	> 7.28%	2.20 (1.12 to 4.31)	Unadjusted HR
Bady 2012 (M-GBM)	Bead array	Frozen	31 and 83	> 0.358	6.46 (2.41 to 17.3)	IPD
	MSP	NR	76–80 and 84–86	NR	7.21 (2.37 to 22.0)	IPD
Barault 2015	Methyl- beaming	FFPE	79–83	> 40.2%	2.78 (1.85 to 5.26)	Unadjusted HR
	PSQ	FFPE	76-81	> 29.6%	2.63 (1.43 to 4.55)	Unadjusted HR
Barbagallo 2014	MSP	FFPE	76–80 and 84–87	Including weakly	3.68 (1.66 to 8.18)	KM curves (Fig 3A or 3C, Barba gallo 2014)
	MSP	FFPE	76–80 and 84–87	Excluding weakly	1.90 (0.72 to 4.99)	KM curves (Fig 3C, Barbagallo 2014)
	PSQ	FFPE	NR	> 9%	3.73 (1.68 to 8.28)	KM curves (Fig 3A or 3B, Barba gallo 2014)
	PSQ	FFPE	NR	> 25%	1.99 (0.92 to 4.32)	KM curves (Fig 3B, Barbagallo 2014)
Bell 2017	QF-IHC (AQUA)	FFPE	N/A	> Median	1.84 (1.38 to 2.43)	Adjusted HR (for age, KPS, re- section status and treatment)
	qMSP	NR	NR	> 8	1.77 (1.28 to 2.44)	Adjusted HR (for age, KPS, re- section status and treatment)
Brigliadori 2016	PSQ	FFPE	74-83	>9%	1.92 (1.17 to 3.14)	KM curves (Fig 1; stratified by extent of resection, Brigliadori 2016)
	PSQ	FFPE	74-83	>29%	3.02 (1.72 to 5.29)	KM curves (Fig 2; stratified by extent of resection, Brigliadori 2016)

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Chai 2018 (7-site co-	PSQ	Frozen	72–78	> 12%	2.94 (1.12 to 7.69)	Unadjusted HR
hort)	PSQ	Frozen	74–78	> 12%	2.94 (1.12 to 7.69)	Unadjusted HR
	PSQ	Frozen	75–78	> 12%	2.94 (1.12 to 7.69)	Unadjusted HR
Chai 2018 (8-site co-	PSQ	Frozen	75–78	> 13%	2.70 (1.37 to 5.26)	Unadjusted HR
hort)	PSQ	Frozen	75-82	> 12%	3.03 (1.54 to 6.25)	Unadjusted HR
	PSQ	Frozen	76–79	> 11%	2.13 (1.09 to 4.17)	Unadjusted HR
Dahlrot 2018 (NS	DIF	FFPE	N/A	< 0.2	1.60 (0.95 to 2.71)	Unadjusted HR
cohort)	PSQ	FFPE	74–78	>9%	1.42 (0.84 to 2.40)	Unadjusted HR
Dahlrot 2018 (RSD	DIF	FFPE	N/A	< 0.2	2.00 (1.32 to 3.02)	Unadjusted HR
cohort)	PSQ	FFPE	74–78	> 10%	1.58 (1.14 to 2.19)	KM curves (Fig 3A, Dahlrot 2018)
Dunn 2009	PSQ	Frozen, smear, FF- PE or a combina- tion	72-83	> 9%	3.57 (2.24 to 5.70)	KM curves (Fig 2B, Dunn 2009)
	PSQ	Frozen, smear, FF- PE or a combina- tion	72-83	> 20%	4.25 (2.57 to 7.05)	KM curves (Fig 2D, Dunn 2009)
	PSQ	Frozen, smear, FF- PE or a combina- tion	72-83	> 29%	4.03 (2.30 to 7.07)	KM curves (Fig 2F, Dunn 2009)
	PSQ	Frozen, smear, FF- PE or a combina- tion	72-83	> 35%	3.64 (1.99 to 6.67)	KM curves (Fig 2D, Dunn 2009)
	PSQ	Frozen, smear, FF- PE or a combina- tion	72–83	Cluster 1 vs 2 and 3	4.44 (2.58 to 7.66)	KM curves (Fig Suppl 4C, Dunn 2009)
	PSQ	Frozen, smear, FF- PE or a combina- tion	72-83	Cluster 1 and 2 vs 3	3.59 (2.26 to 5.69)	KM curves (Fig Suppl 4C, Dunn 2009)
Felsberg 2009	IHC	FFPE	N/A	< 10%	1.26 (0.70 to 2.25)	KM curves (Fig Suppl 1C, 2nd column, Felsberg 2009)

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	MSP	Frozen (14 FFPE)	NR	NR	2.23 (1.29 to 3.84)	KM curves (Fig Suppl 1A, 2nd column, Felsberg 2009)
	PCR-mRNA	Frozen (14 FFPE)	N/A	< 50%	2.66 (0.94 to 7.53)	KM curves (Fig Suppl 1B, 2nd column, Felsberg 2009)
Havik 2012	MSP	Frozen	76–80 and 84–87	NR	2.02 (1.08 to 3.77)	KM curves (Fig 2A, Johannesser 2018)
	PCR-HRM	Frozen	72-83	NR	1.32 (0.64 to 2.68)	KM curves (Fig 2B, Johannesser 2018)
	PSQ	Frozen	74–78	> 2.68%	1.85 (1.02 to 3.35)	KM curves (Fig 1B, Havik 2012)
	PSQ	Frozen	74–78	>6%	2.30 (1.21 to 4.38)	Unadjusted HR
	PSQ	Frozen	74–78	>7%	2.33 (1.19 to 4.57)	Unadjusted HR
	PSQ	Frozen	74–78	> 8%	2.33 (1.19 to 4.57)	Unadjusted HR
	PSQ	Frozen	74–78	>9%	2.30 (0.99 to 5.33)	Unadjusted HR
	PSQ	Frozen	76–79	> 6%	2.30 (1.21 to 4.38)	Unadjusted HR
	PSQ	Frozen	76–79	>7%	2.33 (1.19 to 4.57)	Unadjusted HR
	PSQ	Frozen	76–79	>8%	1.90 (0.99 to 3.65)	Unadjusted HR
	PSQ	Frozen	76–79	>9%	1.90 (0.99 to 3.65)	Unadjusted HR
	qMSP	Frozen	71–73 and 75–86	NR	1.72 (0.91 to 3.22)	KM curves (Fig 2C, Johannesser 2018)
	qMSP	Frozen	71-86	>0%	1.66 (0.97 to 2.83)	KM curves (Fig 1A, Havik 2012)
Hsu 2015	IHC	FFPE	N/A	< 10%	2.12 (1.32 to 3.42)	Unadjusted HR
	MSP	FFPE	76–80 and 84–87	NR	2.39 (1.42 to 4.02)	Unadjusted HR
	PSQ	FFPE	76–79	> 5%	2.66 (1.49 to 4.76)	Unadjusted HR
	qMSP	FFPE	77–80 and 84–87	> 0.04%	2.75 (1.51 to 5.04)	Unadjusted HR
	qMSP	FFPE	77–80 and 84–87	>0.1%	2.83 (1.85 to 4.33)	IPD
Karayan- Tapon 2010	IHC	FFPE	N/A	< 15.5%	1.26 (0.77 to 2.06)	KM curves (Fig 3B, Karayan- Tapon 2010)
	MSP	Frozen	76–80 and 84–87	NR	2.32 (1.39 to 3.87)	KM curves (Fig 1A, Karayan- Tapon 2010)
	PCR-mRNA	Frozen	N/A	< 0.39	1.68 (1.04 to 2.73)	KM curves (Fig 3A, Karayan- Tapon 2010)

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	PSQ	Frozen	74	> 5.5%	3.26 (1.91 to 5.59)	KM curves (Fig Elec Suppl 1B, Karayan-Tapon 2010)
	PSQ	Frozen	74–78	> 8.0%	3.35 (1.95 to 5.73)	KM curves (Fig 2B; cpg mean, Karayan-Tapon 2010)
	PSQ	Frozen	75	> 8.7%	3.26 (1.91 to 5.57)	KM curves (Fig Elec Suppl 1B, Karayan-Tapon 2010)
	PSQ	Frozen	76	> 8.0%	2.78 (1.64 to 4.74)	KM curves (Fig Elec Suppl 1B, Karayan-Tapon 2010)
	PSQ	Frozen	77	> 7.85%	3.65 (2.10 to 6.34)	KM curves (Fig 2B, Karayan- Tapon 2010)
	PSQ	Frozen	78	> 7.8%	2.70 (1.59 to 4.58)	KM curves (Fig Elec Suppl 1B, Karayan-Tapon 2010)
	SQ-MSP	Frozen	76–80 and 84–87	> 35	2.75 (1.66 to 4.53)	KM curves (Fig 1B, Karayan- Tapon 2010)
Kim 2016	MSP	FFPE	76–80 and 84–87	NR	7.66 (2.82 to 20.8)	KM curves (Fig 2G, Kim 2016)
	PSQ	FFPE	74–78	>9%	7.66 (2.82 to 20.8)	KM curves (Fig 2G, Kim 2016)
Kristensen	IHC	FFPE	N/A	At 0%	1.58 (1.12 to 2.22)	Unadjusted HR
2016	PSQ	Frozen	NR	> 10%	1.80 (1.23 to 2.62)	Unadjusted HR
	qMSP-PSQ	Frozen	NR	>0.1%	1.64 (1.15 to 2.33)	Unadjusted HR
	qMSP-PSQ	Frozen	NR	> 5%	1.66 (1.02 to 2.71)	KM curves (Fig 5a, Kristensen 2016)
	qMSP-PSQ	Frozen	NR	> 20%	1.52 (0.77 to 3.00)	KM curves (Fig 5a, Kristensen 2016)
Lalezari	IHC	FFPE	N/A	< 30%	1.74 (1.39 to 2.16)	KM curves (Fig 1A, Lalezari 2013)
2013	MSP	FFPE	76–80 and 84–87	NR	2.13 (1.67 to 2.78)	Adjusted HR (for age, sex, KPS, extent of resection, bevacizum- ab treatment at any time and IDH1R132 mutation status)
	PSQ	FFPE	72–95	NR	2.06 (1.62 to 2.62)	KM curves (Fig 1E; unadjusted, Lalezari 2013)
Lattanzio 2015	MSP	FFPE	76–80 and 84–87	NR	1.45 (0.76 to 2.76)	KM curves (Fig 3B, Lattanzio 2015)
	MSP	Frozen	76–80 and 84–87	NR	2.27 (1.21 to 4.26)	KM curves (Fig 3A, Lattanzio 2015)
	PSQ	FFPE	72-80	≥9%	2.09 (1.09 to 3.99)	KM curves (Fig 3D, Lattanzio 2015)

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(Continued)						
	PSQ	Frozen	72–80	≥9%	2.25 (1.19 to 4.25)	KM curves (Fig 3C, Lattanzio 2015)
Lechapt- Zalcman	IHC	FFPE	N/A	< 15%	1.99 (1.15 to 3.42)	KM curves (Fig 4d; unadjusted, Lechapt-Zalcman 2012)
2012	MSP	FFPE	76–80 and 84–87	NR	1.78 (1.03 to 3.09)	KM curves (Fig 4c; unadjusted, Lechapt-Zalcman 2012)
McDonald 2013	MSP	FFPE	76–80	NR	1.64 (0.95 to 2.83)	IPD
2013	PSQ	FFPE	74-78	> 8%	1.96 (1.16 to 3.33)	Unadjusted HR
Melguizo 2012	IHC	FFPE	N/A	< 25%	1.11 (0.69 to 1.77)	KM curves (Fig 4B, Melguizo 2012)
	MSP	NR	76–80 and 84–87	NR	1.77 (1.06 to 2.95)	KM curves (Fig 4A, Melguizo 2012)
Nguyen 2015	FSQ-MS- PCR	Frozen or FFPE	76–80 and 84–87	> 15%	2.68 (1.70 to 4.21)	KM curves (Fig 3D, Nguyen 2015)
	FSQ-MS- PCR	Frozen or FFPE	76–80 and 84–87	> 60%	2.25 (1.31 to 3.87)	KM curves (Fig 3D, Nguyen 2015)
Park 2011	MS-MLPA	50% frozen and 50% FFPE	NR	> 0.1%	2.38 (1.11 to 5.10)	KM curves (Fig 2B, Park 2011)
	MS-MLPA	50% frozen and 50% FFPE	NR	> 0.2	1.88 (0.86 to 4.11)	KM curves (Fig 2B, Park 2011)
	MSP	50% frozen and 50% FFPE	76–80 and 84–86	NR	4.53 (1.58 to 12.9)	KM curves (Fig 2A, Park 2011)
Quillien 2014 (test)	IHC	FFPE	N/A	< 23%	2.33 (1.44 to 3.74)	Adjusted HR (for age and KPS)
2014 (lest)	MSP	Frozen	76–80 and 84–87	NR	2.70 (1.65 to 4.43)	Adjusted HR (for age and KPS)
	Methy- Light-MSP	Frozen	75–86	> 0	1.67 (1.00 to 2.77)	Adjusted HR (for age and KPS)
	PCR-HRM	Frozen	70-83	> 50%	1.92 (1.12 to 3.29)	Adjusted HR (for age and KPS)
	PSQ	Frozen	74	> 4%	2.44 (1.51 to 3.95)	Adjusted HR (for age and KPS)
	PSQ	Frozen	74	> 8%	2.22 (1.30 to 3.79)	Adjusted HR (for age and KPS)
	PSQ	Frozen	74–78	> 8%	3.13 (1.86 to 5.25)	Adjusted HR (for age and KPS)
	PSQ	Frozen	74–78	> 9%	3.13 (1.81 to 5.38)	Adjusted HR (for age and KPS)
	PSQ	Frozen	74-89	>11%	3.13 (1.84 to 5.30)	Adjusted HR (for age and KPS)

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	PSQ	Frozen	75	> 11%	3.23 (1.87 to 5.55)	Adjusted HR (for age and KPS)
	PSQ	Frozen	75–79	> 8%	2.94 (1.71 to 5.06)	Adjusted HR (for age and KPS)
	PSQ	Frozen	76	> 4%	2.63 (1.61 to 4.31)	Adjusted HR (for age and KPS)
	PSQ	Frozen	76	> 5%	2.78 (1.62 to 4.75)	Adjusted HR (for age and KPS)
	PSQ	Frozen	76–79	>8%	2.86 (1.66 to 4.92)	Adjusted HR (for age and KPS)
	PSQ	Frozen	76-80	>9%	3.03 (1.74 to 5.26)	Adjusted HR (for age and KPS)
	PSQ	Frozen	77	> 6%	3.13 (1.87 to 5.23)	Adjusted HR (for age and KPS)
	PSQ	Frozen	77	>7%	2.78 (1.63 to 4.74)	Adjusted HR (for age and KPS)
	PSQ	Frozen	77-81	> 8%	2.70 (1.59 to 4.59)	Adjusted HR (for age and KPS)
	PSQ	Frozen	78	> 4%	2.33 (1.38 to 3.92)	Adjusted HR (for age and KPS)
	PSQ	Frozen	78	> 5%	2.50 (1.51 to 4.13)	Adjusted HR (for age and KPS)
	PSQ	Frozen	78-82	>9%	2.86 (1.68 to 4.86)	Adjusted HR (for age and KPS)
	PSQ	Frozen	79	>7%	2.78 (1.61 to 4.80)	Adjusted HR (for age and KPS)
	PSQ	Frozen	79-83	>8%	2.86 (1.68 to 4.86)	Adjusted HR (for age and KPS)
	PSQ	Frozen	80	> 4%	2.56 (1.54 to 4.28)	Adjusted HR (for age and KPS)
	PSQ	Frozen	81	>8%	2.44 (1.44 to 4.14)	Adjusted HR (for age and KPS)
	PSQ	Frozen	82	> 16%	2.94 (1.70 to 5.08)	Adjusted HR (for age and KPS)
	PSQ	Frozen	83	>10%	2.78 (1.65 to 4.66)	Adjusted HR (for age and KPS)
	PSQ	Frozen	84	>9%	3.23 (1.86 to 5.59)	Adjusted HR (for age and KPS)
	PSQ	Frozen	84-88	> 17%	3.23 (1.86 to 5.58)	Adjusted HR (for age and KPS)
	PSQ	Frozen	84-89	> 22%	3.23 (1.85 to 5.62)	Adjusted HR (for age and KPS)
	PSQ	Frozen	85	> 5%	2.50 (1.52 to 4.11)	Adjusted HR (for age and KPS)
	PSQ	Frozen	85-89	> 13%	2.94 (1.76 to 4.92)	Adjusted HR (for age and KPS)
	PSQ	Frozen	86	> 11%	2.78 (1.65 to 4.69)	Adjusted HR (for age and KPS)
	PSQ	Frozen	87	> 25%	3.03 (1.75 to 5.24)	Adjusted HR (for age and KPS)
	PSQ	Frozen	88	>4%	2.27 (1.38 to 3.75)	Adjusted HR (for age and KPS)
	PSQ	Frozen	89	> 12%	3.23 (1.91 to 5.46)	Adjusted HR (for age and KPS)
Quillien 2014 (vali-	PSQ	FFPE	74–78	> 9%	3.70 (1.71 to 8.01)	Unadjusted HR

dation)

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Continued)						
	PSQ	FFPE	74–78	> 10%	2.86 (1.42 to 5.74)	Unadjusted HR
	PSQ	FFPE	74–78	> 28%	2.27 (0.98 to 5.29)	Unadjusted HR
Quillien 2016	PSQ	FFPE	74–78	> 6%	3.23 (2.02 to 5.16)	Adjusted HR (for age and KPS)
2010	PSQ	FFPE	74–78	> 8%	4.00 (2.30 to 6.97)	Adjusted HR (for age and KPS)
	PSQ	FFPE	74–78	> 12%	4.17 (2.35 to 7.38)	Adjusted HR (for age and KPS)
	PSQ	FFPE	74–78	> 13%	4.35 (2.41 to 7.83)	Adjusted HR (for age and KPS)
	PSQ	FFPE	74–78	> 16%	4.55 (2.48 to 8.34)	Adjusted HR (for age and KPS)
	PSQ	Frozen	74–78	> 6%	4.00 (2.30 to 6.97)	Adjusted HR (for age and KPS)
	PSQ	Frozen	74–78	> 8%	3.57 (2.14 to 5.95)	Adjusted HR (for age and KPS)
	PSQ	Frozen	74–78	> 12% or 13%	3.45 (2.10 to 5.66)	Adjusted HR (for age and KPS)
	PSQ	Frozen	74–78	> 12%	3.70 (2.19 to 6.26)	Adjusted HR (for age and KPS)
	PSQ	Frozen	74–78	> 16%	3.13 (1.94 to 5.04)	Adjusted HR (for age and KPS)
	PSQ	Frozen	76–79	> 8%	3.33 (2.06 to 5.40)	Adjusted HR (for age and KPS)
	PSQ	Frozen	76–79	> 12%	3.33 (2.06 to 5.40)	Adjusted HR (for age and KPS)
	SQ-MSP	FFPE	76–80 and 84–87	> 12%	3.33 (2.06 to 5.40)	Adjusted HR (for age and KPS)
	SQ-MSP	FFPE	76–80 and 84–87	> 13%	3.33 (2.06 to 5.40)	Adjusted HR (for age and KPS)
	SQ-MSP	FFPE	76–80 and 84–87	> 23%	4.17 (2.35 to 7.38)	Adjusted HR (for age and KPS)
	SQ-MSP	Frozen	76–80 and 84–87	> 13%	2.86 (1.79 to 4.57)	Adjusted HR (for age and KPS)
	SQ-MSP	Frozen	76–80 and 84–87	> 23%	2.17 (1.38 to 3.44)	Adjusted HR (for age and KPS)
Thon 2017	MSP	Frozen	76–80 and 84–87	NR	3.33 (1.82 to 6.25)	Unadjusted HR
	Sequencing	Frozen	75–99 (un- clear)	> 50%	3.33 (1.82 to 6.25)	Unadjusted HR
Yamashita 2018	MSP	Frozen	76–80 and 84–87	NR	2.36 (1.62 to 5.05)	Unadjusted HR
	PCR-HRM	Frozen	72-89	> 5%	2.36 (1.43 to 3.90)	KM curves (Fig Suppl 13, Ya- mashita 2018)

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(Continued)						
	PCR-HRM	Frozen	72–89	> 8%	2.05 (1.24 to 3.39)	KM curves (Fig Suppl 13, Ya- mashita 2018)
	PCR-HRM	Frozen	72–89	> 10%	2.51 (1.63 to 4.83)	Unadjusted HR
	PCR-HRM	Frozen	72–89	> 12%	2.61 (1.54 to 4.42)	KM curves (Fig Suppl 13, Ya- mashita 2018)
	PCR-HRM	Frozen	72–89	> 15%	2.39 (1.41 to 4.05)	KM curves (Fig Suppl 13, Ya- mashita 2018)
Yang 2012	IHC	FFPE	N/A	< 10%	1.07 (0.35 to 3.31)	IPD
	MSP	FFPE	76–80 and 84–87	NR	1.35 (0.44 to 4.16)	IPD
Yoshioka 2018	SQ-MSP	Frozen	76–80 and 84–87	> 0	2.72 (1.28 to 5.74)	KM curves (Fig 2A, Yoshioka 2018)
	SQ-MSP	Frozen	76–80 and 84–87	>2	2.18 (1.20 to 3.97)	KM curves (Fig 2B, Yoshioka 2018)
	SQ-MSP	Frozen	76–80 and 84–87	>4	1.85 (1.07 to 3.18)	KM curves (Fig 2C, Yoshioka 2018)
	SQ-MSP	Frozen	76–80 and 84–87	>6	1.83 (1.10 to 3.04)	KM curves (Fig 2D, Yoshioka 2018)
	SQ-MSP	Frozen	76–80 and 84–87	> 8	1.71 (1.00 to 2.93)	KM curves (Fig 2E, Yoshioka 2018)

CpG: 5'-cytosine-phosphate-guanine-3'; DIF: double immunofluorescence; FFPE: formalin-fixed paraffin-embedded; FSQ-MS-PCR: fluorescent semi-quantitative methylation-specific polymerase chain reaction; HR: hazard ratio; IDH: isocitrate dehydrogenase; IHC: immunohistochemistry; IPD: individual participant data; KM: Kaplan-Meier; KPS: Karnofsky performance status; MS-MLPA: methy-lation-specific multiplex ligation-dependent probe amplification; MS-RE-qPCR: methylation-specific restriction enzyme quantita-tive polymerase chain reaction; MSP: methylation-specific polymerase chain reaction; N/A: not applicable; NR: not reported; PCR: polymerase chain reaction; PCR-HRM: polymerase chain reaction with high-resolution melting; PCR-mRNA: polymerase chain reaction-messenger ribonucleic acid; PSQ: pyrosequencing; QF-IHC: quantitative fluorescence immunohistochemistry; qMSP: quantitative methylation-specific polymerase chain reaction with pyrosequencing; SQ-MSP: semi-quantitative methylation-specific polymerase chain reaction with pyrosequencing; SQ-MSP: semi-quantitative methylation-specific polymerase chain reaction.

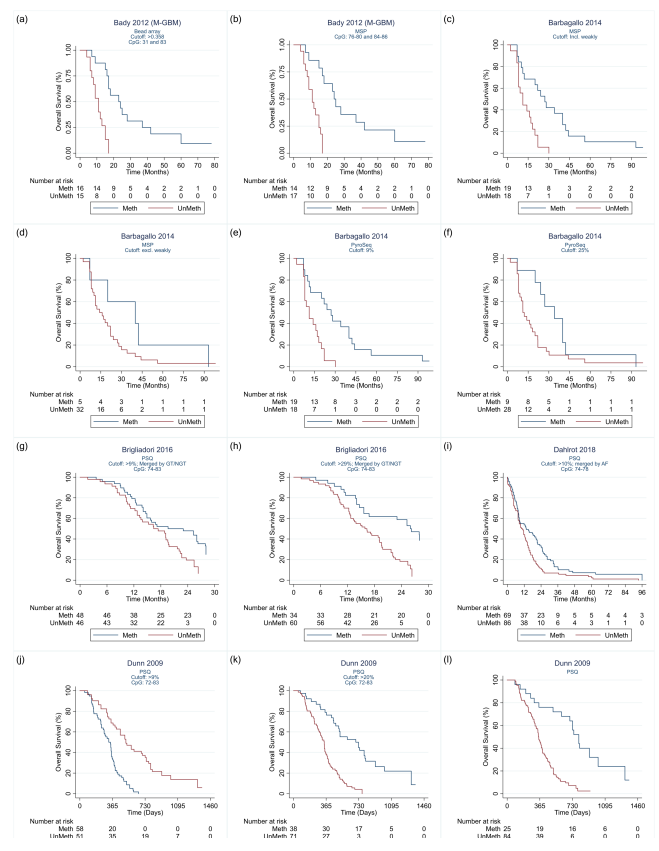
Appendix 9. Reconstructed Kaplan-Meier plots

Reconstructed Kaplan-Meier plots based on reported IPD or published Kaplan-Meier curves are presented in Figure 16, Figure 17, Figure 18, Figure 19, Figure 20, and Figure 21.

Figure 16. Reconstructed Kaplan-Meier curves (1/6). AF: area fraction; CpG: 5'-cytosine-phosphate-guanine-3'; GT: gross total; Incl: including; M-GBM: methylated glioblastoma; Meth: methylated; MSP: methylation-specific

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polymerase chain reaction; NGT: non-gross total; PSQ: pyrosequencing; PyroSeq: pyrosequencing; UnMeth: unmethylated.



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Figure 16. (Continued)

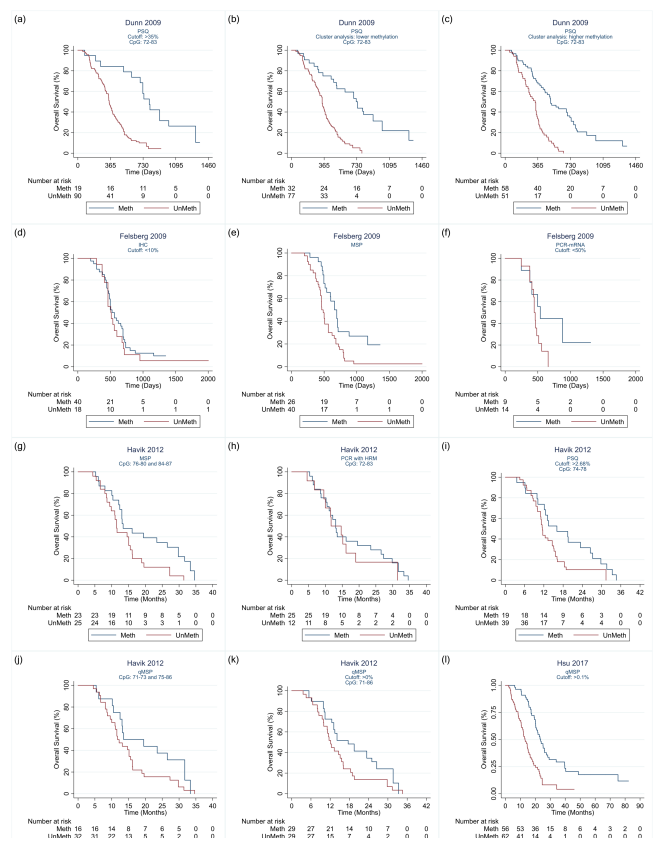
		lime (Days)				i ime (Days)	Lime (Days)							
Number at risk					Number at risk					Number at risk						
Meth 58	20	0	0	0	Meth 38	30	17	5	0	Meth 25	19	16	6	0		
UnMeth 51	35	19	7	0	UnMeth 71	27	3	0	0	UnMeth 84	39	6	0	0		
	Me	eth ——	 UnMet 	۱	-	Me	eth —	- UnMet	h	-	Me	eth ——	 UnMeti 	n		

Figure 17. Reconstructed Kaplan-Meier curves (2/6). CpG: 5'-cytosine-phosphate-guanine-3'; IHC: immunohistochemistry; Meth: methylated; MSP: methylation-specific polymerase chain reaction; NGT: ; PCR-HRM: polymerase chain reaction with high-resolution melting; PCR-mRNA: polymerase chain reaction targeting

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messenger ribonucleic acid; PSQ: pyrosequencing; qMSP: quantitative methylation-specific polymerase chain reaction; UnMeth: unmethylated.



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Figure 17. (Continued)

			Lime	e (Moi	ntns)							ıme (N	nontns	,						Lin	ne (N	/iontn	s)			
Number at risk									Number at risk								Number at risk									
Meth 16	16	14	8	7	6	5	0	0	Meth 29	27	21	14	10	7	0	0	Meth 56	53	36	15	8	6	4	3	2	0
UnMeth 32	31	22	13	5	5	2	0	0	UnMeth 29	27	15	7	4	2	0	0	UnMeth 62	41	14	4	1	0	0	0	0	0
			Meth	_		- Uni	/leth				— м	eth		— Un	Meth					Met	h		_	UnM	eth	

Figure 18. Reconstructed Kaplan-Meier curves (3/6). CpG: 5'-cytosine-phosphate-guanine-3'; IHC: immunohistochemistry; Meth: methylated; MSP: methylation-specific polymerase chain reaction; NA: not

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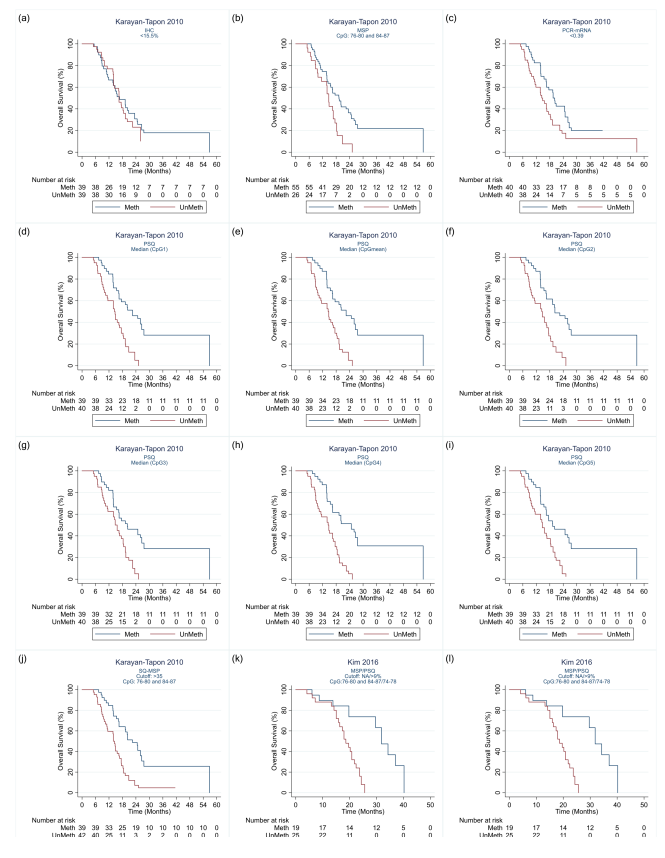
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available; PCR-mRNA: polymerase chain reaction-messenger ribonucleic acid; PSQ: pyrosequencing; SQ-MSP: semiquantitative methylation-specific polymerase chain reaction; UnMeth: unmethylated.



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Figure 18. (Continued)

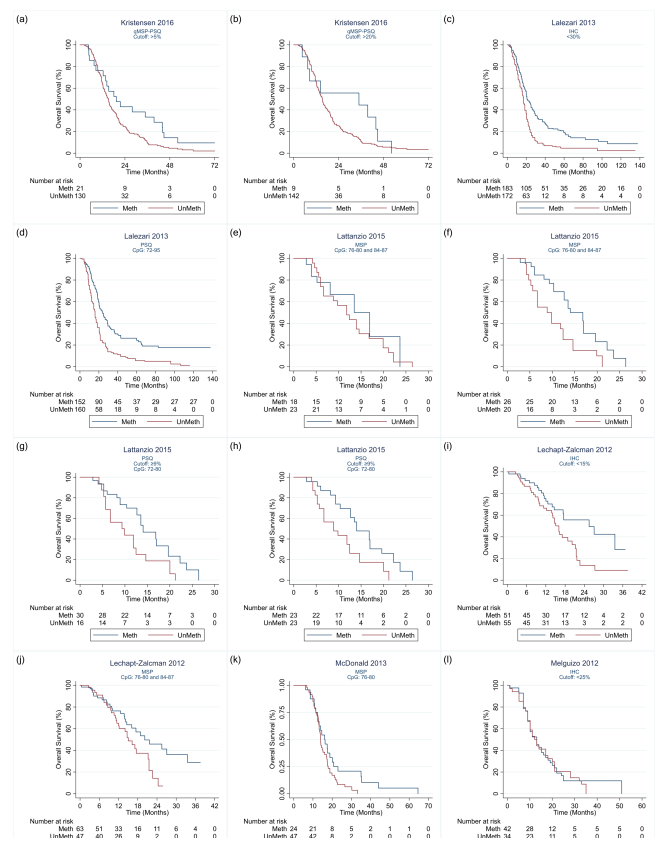
lime (Months)				I ime (I	viontns)				I ime (viontns)			
Number at risk		Number at risk						Number at risk					
Meth 39 39 33 25 19 10 10 10	10 10 0	Meth 19	17	14	12	5	0	Meth 19	17	14	12	5	0
UnMeth 42 40 25 11 3 2 2 0	0 0 0	UnMeth 25	22	11	0	0	0	UnMeth 25	22	11	0	0	0
Meth U	Meth	[—— I	Meth		UnMeth		[Meth		UnMeth	

Figure 19. Reconstructed Kaplan-Meier curves (4/6). CpG: 5'-cytosine-phosphate-guanine-3'; IHC: immunohistochemistry; Meth: methylated; MSP: methylation-specific polymerase chain reaction; PSQ:

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pyrosequencing; qMSP-PSQ: quantitative methylation-specific polymerase chain reaction with pyrosequencing; UnMeth: unmethylated.



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Figure 19. (Continued)

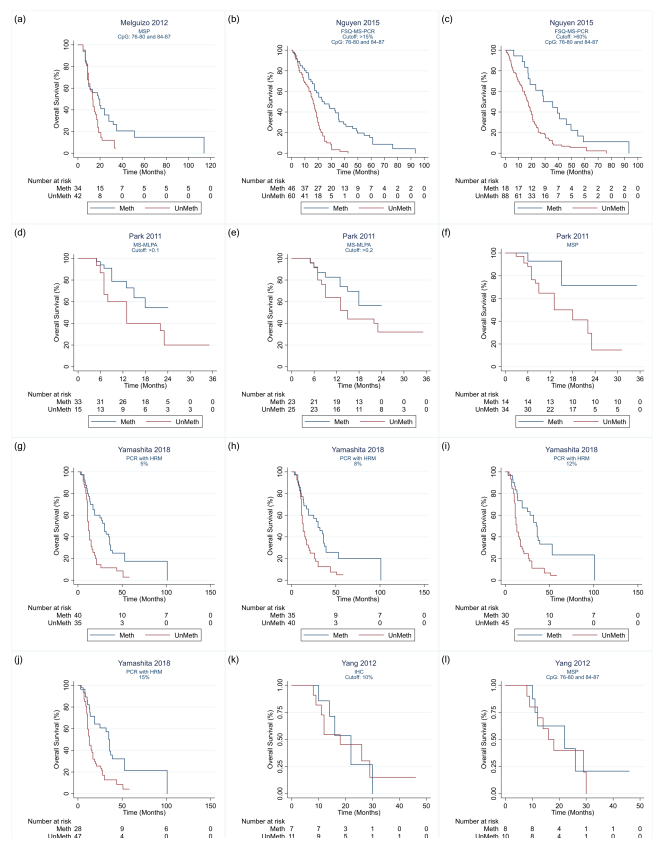
		1	ıme (r	nontns)						ıme (r	nontns)					LIM	e (Mon	tns)		
Number at risk Meth 63	51	33	16	11	6	4	0	Number at risk Meth 24	21	8	5	2	1	1	0	Number at risk Meth 42	28	12	5	5	5	0
UnMeth 47	40	26	9	2	Ő	ò	Ő	UnMeth 47	42	8	2	ō	ò	0	Ő	UnMeth 34	23	11	5	Ő	õ	õ
Meth UnMeth										— м	leth		— Ur	Meth				 Meth 		ι	JnMeth	

Figure 20. Reconstructed Kaplan-Meier curves (5/6). CpG: 5'-cytosine-phosphate-guanine-3'; FSQ-MS-PCR: fluorescent semi-quantitative methylation-specific polymerase chain reaction; IHC: immunohistochemistry; Meth: methylated; MS-MLPA: methylation-specific multiplex ligation-dependent probe amplification; MSP: methylation-

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specific polymerase chain reaction; PCR with HRM: polymerase chain reaction with HRM: high-resolution melting; UnMeth: unmethylated.



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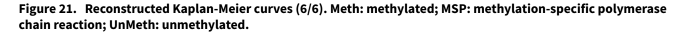
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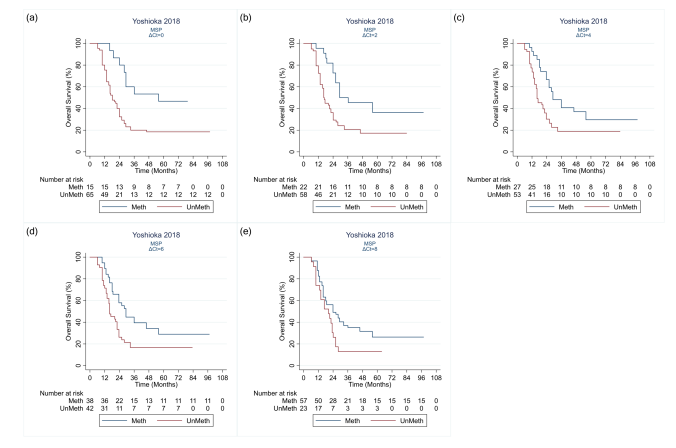
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Figure 20. (Continued)

	Lime	(Months)				Iime	(Months)				Iime	(Months)			
Number at risk Meth 28	9	6	0	Number at risk Meth 7	7	3	1	0	0	Number at risk Meth 8	8	4	1	1	0
UnMeth 47	4	Ő	õ	UnMeth 11	9	5	1	1	õ	UnMeth 10	8	4	<u>i</u>	Ö	ŏ
	Meth	UnMeth				Meth		- UnMeth				Meth		- UnMeth	1





HISTORY

Protocol first published: Issue 4, 2019 Review first published: Issue 3, 2021

CONTRIBUTIONS OF AUTHORS

AM led development of the protocol and organised retrieval of papers. SD designed and undertook the searches. AM, KMK and AH screened search results. AM, CK, FS, KMK, AH, SB and CLF screened retrieved papers against eligibility criteria. AK and TR screened retrieved papers against economic eligibility criteria. AM, CK, FS, LS, HC and JPTH extracted data from papers. AM, CK, FS, LS, HC and JPTH extracted data from papers. AM, CK, FS, LS, HC and JPTH assessed risk of bias and applicability of included studies. JPTH and HC undertook the statistical analyses. JPTH undertook GRADE assessments. KMK, JPTH, FS and CLF interpreted the results. JPTH and KMK co-ordinated the review. KMK, CLF, SB, SJ and CW provided a clinical perspective. LV, AK and TR provided economics expertise.

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JPTH drafted the manuscript with contributions from KMK, FS and HC.

DECLARATIONS OF INTEREST

AM: none known CK: none known FS: none known AK: none known HC: none known SD: none known LS: none known TR: none known SB: member of the NICE Primary Brain Tumours Guideline Committee CLF: none known CW: none known SJ: member of the National Institute for Health and Care Excellence (NICE) Primary Brain Tumours Guideline Committee AH: none known LV: none known JPTH: none known KMK: none known

SOURCES OF SUPPORT

Internal sources

• University of Bristol, UK

External sources

• National Institute for Health Research (NIHR), UK

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• Cancer Research UK, UK

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The views and opinions expressed are those of the review authors and do not necessarily reflect those of the NIHR, its Systematic Reviews Programme, the National Health Service (NHS), the Department of Health or Cancer Research UK.

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We had planned to search Open Grey (www.opengrey.eu/) using the free text terms from our MEDLINE search and in dissertations and theses using ProQuest Dissertations & Theses Global (search.proquest.com/pqdtglobal/dissertations/) and Networked Digital Library of Theses and Dissertations (search.ndltd.org/index.php). We did not implement these.

We had planned meta-analyses (if deemed appropriate), although had not clarified that these would target *comparisons* of techniques (or of variants of techniques) to address the prespecified objectives of the review. Our meta-analyses all addressed such comparisons. We developed our specific approach for this while undertaking the analyses (specifically, our imputation of a correlation between log HRs based on correlations among tests results in some individual participant data that we found in publications of included studies).

We rearranged the domains of our risk of bias assessment, although it covers the same issues and uses the same signalling questions. This was to allow us to separate better the study-level issues from the result-level issues.

We had aimed to create an economic model using outcomes from both the clinical and economic evidence. This was to be a decision analytic model, assessing the cost-effectiveness of different methods of testing for MGMT promoter methylation status in people with glioma. We decided against this, in consultation with the hosting Cochrane Review Group, on the basis that it would involve a comparison of different policies for managing people using the information about MGMT promoter methylation status, thus extending substantially beyond the scope of the review.

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