Brain metastases (BM) in patients with non-small cell lung cancer (NSCLC)

Despite the gradual treatment improvements of primary sites of NSCLC, still one of the biggest challenges is how to prevent or treat BM successfully in this group. In about 15% of patients with NSCLC, the brain will be a site of first recurrence (1). At diagnosis 10–20% of patients with NSCLC present BM. In stage I approximately 30% of patients will develop BM during the initial 2 years since diagnosis with the figure reaching 40–50% for the whole duration of treatment irrespectively of surgery or diagnostic tools used.

The current standard for diagnosing BM is MRI, which is recommended for patients with stage II or higher (and optionally used in Ib) according to NCCN guidelines v7.2019 (2). Should MRI be impossible, CT with contrast is suggested as an alternative regardless of the presence
of any neurological symptoms. BM have a devastating impact on the patients’ overall survival (OS), with a median of 6 months of survival after the manifestation of BM, and severely impair the quality of life (QoL). Presence of specific mutations in the tumor may considerably alter the course of the disease. Patients with EGFR mutations or ALK rearrangements are less prone to worse prognosis. Patients presenting Diagnosis Specific Graded Prognostic Index (DS_GPA) assessment/Assessment for Lung Cancer Using Molecular Markers (Lung-mol GPA) scores 3.5–4.0 may have a median of survival of 4 years (3). This also implicates the necessity to protect the abovementioned group of patients from life threatening aspects of the disease, including BM occurrence. To do so efficiently, one has to identify the groups that may benefit the most from interventions to prevent BM, discuss the options with the patients and consider the costs of such treatment.

**Prevention of BM**

Use of methods aimed at preventing BM should be considered despite the apparent lack of OS benefit as such treatment may preserve adequate neurologic and psychological aspects of QoL (4,5). Higher BM rates are also associated with a greater extent of disease and as such, as evidenced by studies such as RT0G 0214 countering BMs could also be considered as means to prolong disease free survival (DFS) itself (6). It is well known that BM themselves may deteriorate neurological and physical function and thus becoming symptomatic BM. There are several options, which are to be considered as potential BM prevention tools.

**Chemotherapy, molecular targets, immune checkpoint inhibitors used as a prevention tool for BM occurrence**

Chemotherapy currently in routine use or used experimentally does not seem to efficiently protect patients from BM (7). The reason is the function of the blood-brain barrier (BBB), which prohibits cytotoxic agents from reaching brain tissue by adequate concentrations. Data, so far accumulated only in non-squamous NSCLC, shows that pemetrexed used as first line therapy in advanced NSCLC may reduce the risk of developing BM as the first site of progression (8). These observations are however derived from a retrospective analysis of two randomized trials and significant results were obtained only in subgroup analyses limiting their level of evidence. Therefore, this observation has to be put into more rigorous evaluation to ascertain the real association between usage of pemetrexed and decrease the risk of BM. There were also reports involving temozolomide usage after surgery or after platinum-based chemotherapy as a preventative measure for BM. No significant associations in terms of incidence reduction of BM in either of the cohorts were noted (9). However, there are promises data regarding patients with NSCLC stage III suggesting that durvalumab administered after chemotherapy may decrease the incidence of BM (10). In the PACIFIC study the BM incidence was reduced 2-fold: 5.5% in the durvalumab arm vs. 11% in the control group. Durvalumab seemed to be also effective in an early analysis of a phase II trial with existing non-symptomatic, small BM, in patients with PD-L1 positive melanoma or NSCLC (11). The numbers available were, however, very low: 18 patients with NSCLC and 18 with melanoma limited the weight of this piece of evidence. Nevertheless, the responses to treatment were durable in all patients but one with NSCLC. The safety profile of this study in NSCLC patients showed grade 3 non-neurological complications such as colitis, pneumonitis fatigue or hyperkalemia to occur in some individuals alas no neurologic toxicity of grade 3 or higher was noted (11).

There are also promising initial reports of studies with osimertinib (12) and alectinib (13) aimed at preventing BM. Alectinib is a representant of modern generation of ALK inhibitors that seems to show better penetration through the BBB than the first generation of ALK inhibitors such as crizotinib, for which BBB concentrations equaled only 50% of blood concentrations (14). The efficacy of alectinib in terms of BM risk reduction was studied both in patients with or without previous brain irradiation (13,15) and also in a comparative study with crizotinib (16). Alectinib vs. crizotinib was well studied in the Japanese population in J-ALEX study. In this trial the alectinib arm presented an advantage vs. crizotinib in terms of BM progression. It is of note that in this trial the proportion of patients with BM was higher in the crizotinib group than in the alectinib-treated one (28% vs. 14%, respectively). In a subgroup analysis both patients with BM and no BM gained advantage of a prolonged progression free survival (PFS) in favor to alectinib (16). According to authors of aforementioned study—only patients without BM or with BM and no previous brain radiotherapy gained advantage in terms of PFS. The authors claim that the safety profile of alectinib was better than that of crizotinib, but only non-
The efficacy of osimertinib, a third generation of ALK inhibitors, was assessed in the FLAURA trial (12,17). This randomized, double blind phase III study included Asian and white population in a balanced manner and in nearly half of each arm (osimertinib vs erlotinib or gefitinib) patients presented BM. In the experimental arm osimertinib reduced the frequency of BM progression or death caused by BM down to 52%. Safety profile was significantly in favor of osimertinib in contrast to erlotinib or gefitinib used in the control arm. Non-neurologic adverse events of grade 3 or higher occurred in 34% vs. 45% of patients, respectively. Fatal adverse events occurred in 6 patients (2%) in the osimertinib group and in 10 patients in the standard EGFR-TKI group, but none of them were directly related to osimertinib use (12).

**Prophylactic cranial irradiation as a preventative tool for BM**

Given that in small-cell lung cancer (SCLC) patients prophylactic cranial irradiation (PCI) lead to the reduction of the incidence of BM and prolonged OS (18) further investigations were made in NSCLC patients. So far, PCI significantly reduces the incidence of BM and prolongs the disease-free interval in patients with NSCLC, but thus far no improvements of OS were noted (19,20) even in long term observations. No randomized trials reported significant improvements of OS after PCI in patients with NSCLC. In none of the subgroups, OS prolongation was noted if PCI was used in those cohorts. This undermined the impact of PCI on the prolongation of OS regardless of patients’ stage (21). The lack of impact on OS may result from a variety of causes including but not limited to the majority of patients developing extracranial metastases, low statistical power of recent studies or a true lack of biological effect. The absent effect in terms of OS prolongation may be also the result of slower accrual to studies than initially planned. It should be also taken into consideration that the treatment of BM themselves used in the control arms of studies (i.e., stereotactic radiosurgery, palliative whole brain radiotherapy, steroid intake) does seem to have a beneficial effect, showing longer than average OS times in patients not participating in such clinical trials.

Nonetheless, PCI exerts its beneficial effect by reducing the occurrence of BM by 50–70% (starting from 13% incidence in non-PCI arms with reduction by 6%) (21,22). In a recent publication of De Ruyscher et al., a meta-analysis was performed and showed that the risk ratio of developing BM when PCI is used was 0.33 (95% CI, 0.24–0.45) (1). This effect was significant despite different radiotherapy schemes used in the trials included in the metaanalysis. It is also noteworthy that the most popular radiotherapy protocols used were 30 Gy given in 10 fractions/daily 5 days a week or 30 Gy in 12 fractions in 5 days a week (see Table 1).

At least seven trials, starting from middle late 70-ies through middle 80s (23-26) and recent millennial trials presented a beneficial impact of BM in terms of reduced incidence of BM with the quality of this evidence rated as moderate by a recent systematic review (21). According to systematic reviews and several forms of bias assessment in such papers, despite nearly 40 years of performing such trials the data are still subject to a debate. In terms of that in one systematic review, some works highlight the particular risk of conducting bias (28). In those reviewers’ opinion, however, it was finally assessed as low risk of introducing bias. The latest studies showed that the incidence of BM was in fact higher than expected in both arms, probably as a consequence of better diagnostic tools. In studies performed in late 70s–80s only limited image brain diagnostics were available and were not always performed prior to study enrolment. In studies, which have started in the new millennium, the inclusion criteria involved baseline brain diagnostics performed with at least a CT with contrast evaluation but preferably using high-quality MRI. Table 1 contains an outline of the main studies which included PCI as part of treatment of NSCLC. Trials from the XXth century (apart from SWOG study) recruited patients with diseases of stage I to III. In recent randomized multicenter studies (NVALT11/DLCRG-02 or RTOG 0214, Li et al.), including those with long-term observations, the focus is on patients with locally advanced diseases (stage IIIA–IIB–AJCC 7th edition), who are expected to be at a greater risk of developing BM. In the above listed studies BM were considered as a primary endpoint or secondary one after OS or DFS. In the NVALT11/DLCRG-02 study no significant differences were shown in terms of DFS at 2 years: 67% in PCI arm vs. 72% in the observation arm (1). On the other hand, the long-term observation of in RTOG 0214 showed a significantly prolonged DFS in patients receiving PCI (6). In the Li et al. study, where patients of Asian ethnicity were recruited, the DFS was used as a primary endpoint and was significantly longer in the PCI than in the control arms showing a difference of median survival times of 28.5 vs. 21.2 months (27).
<table>
<thead>
<tr>
<th>Study</th>
<th>Year of study</th>
<th>PCI dose/scheme</th>
<th>Interventions</th>
<th>Toxixity/Qol/neuropsychological tests</th>
<th>Median survival (PCI vs. control)</th>
<th>BM events (PCI vs. control)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cox et al. (VALG) (23)</td>
<td>1981</td>
<td>20 Gy (2×10 Gy)</td>
<td>Arm 1: intermediate-course chest RT (50 Gy/25F/5 weeks); Arm 2: intermediate course chest RT (50 Gy/25F/5 weeks) + PCI; Arm 3: short-course chest RT (42 Gy/15F/3 weeks); Arm 4: short-course chest RT (42 Gy/15F/3 weeks) + PCI</td>
<td>No</td>
<td>8.2 vs. 9.7 months; P=0.5</td>
<td>6% vs. 16%; P=0.038</td>
</tr>
<tr>
<td>Umsawasdi et al. (MDACC) (24)</td>
<td>1984</td>
<td>30 Gy (3×10 Gy)</td>
<td>PCI vs. no-PCI</td>
<td>No Qol</td>
<td>22% vs. 23.5% (at 3 years), P not reported</td>
<td>2% vs. 14%; P=0.002</td>
</tr>
<tr>
<td>Russell et al. (RTOG8403) (25)</td>
<td>1991</td>
<td>30 Gy (3×10 Gy)</td>
<td>PCI given concurrently with the sixth fraction of chest irradiation (30 Gy/10F/2 weeks) primary thoracic RT (55–60 Gy/30F/6 weeks) with no concurrent chemotherapy, while the remaining received postoperative RT (50 Gy/25F/5 weeks) after gross intrathoracic disease resection</td>
<td>No</td>
<td>8.4 vs. 8.1 months; P=0.36</td>
<td>8% vs. 18%; P=0.36</td>
</tr>
<tr>
<td>Mira et al. (SWOG) (26)</td>
<td>1998</td>
<td>30 Gy (2×15 Gy)</td>
<td>Arm 1: chest RT alone (58 Gy/29F/6 weeks); Arm 2: chest RT + PCI; Arm 3: chest RT + CT; Arm 4: chest RT + CT + PCI</td>
<td>No Qol</td>
<td>8 vs. 11 months; P=0.004</td>
<td>1% vs. 13%; P=0.003</td>
</tr>
<tr>
<td>Gore et al. (RTOG 0214 primary results) (20)</td>
<td>2011</td>
<td>30 Gy (2×15 Gy)</td>
<td>High-dose chest irradiation (RT; &gt;30 Gy) with or without adjuvant or neoadjuvant, chemotherapy and/or surgical resection wherever applicable. RT with or without chemotherapy could be given pre- or post-operatively in surgical candidates</td>
<td>NCF and Qol</td>
<td>75.6% vs. 76.9% (1 years); P=0.057</td>
<td>19% vs. 39%; P=0.009</td>
</tr>
<tr>
<td>Li et al. (27)</td>
<td>2015</td>
<td>30 Gy (10×3 Gy)</td>
<td>PCI vs. no-PCI; surgical resection followed by adjuvant platinum-based chemotherapy</td>
<td>Qol (FACT-L) and CTC-AE; RTOG/EORTCLRMSS</td>
<td>44.5% vs. 38.7% (3 years); 27.4% vs. 22.8% (5 years); P=0.31</td>
<td>12.3% vs. 38.7%; P&lt;0.001</td>
</tr>
<tr>
<td>Sun et al. (RTOG 0214 recent results) (6)</td>
<td>2019</td>
<td>30 Gy (15×2 Gy)</td>
<td>PCI vs. no-PCI; RT alone; Chemo-RT</td>
<td>MMSE, ADL, HVLT, acute and late TOX; EORTC QLQ-C30 and BN-20</td>
<td>25.8 vs. 24.8 months; P=0.86</td>
<td>8% vs. 18.1%; P=0.004</td>
</tr>
<tr>
<td>De Ruyscher et al. (NAFALT 11/ DLCRG02) (1)</td>
<td>2018</td>
<td>36 (12× 3 Gy); 30 Gy (12× 2.5 Gy)</td>
<td>PCI vs. no-PCI; RT alone</td>
<td>Qol (EORTC QLQ-C30, BN20, EuroQol 5D), TOX, Neurological events (memory impairment, CTC-AE)</td>
<td>24.2 vs. 21.9 months; P=0.56</td>
<td>7% vs. 27.2%; P&lt;0.001</td>
</tr>
</tbody>
</table>

PCI, prophylactic cranial irradiation; OS, overall survival; NSCLC, non-small cell lung cancer; Gy, Gray; F, number of fractions; RT, radiotherapy; Qol, quality of life; TOX, toxicity; CTC-AE, Common Toxicity Criteria - Adverse Events; EORTC, European Organization for Research and Treatment of Cancer; RTOG/EORTC-LRMSS, RTOG/EORTC late radiation morbidity scoring schema; EORTC QLQ-C30, EORTC quality of life questionnaire-C30; EORTC QLQ-BN20, EORTC quality of life questionnaire-BN20; FACT-L questionnaire, Functional Assessment of Cancer Therapy-Lung questionnaire.
important aspect of this study was that all patients analyzed in that paper had undergone surgery—which may have been a favorable factor pro allowing the authors to obtain adequately long time of observation.

In all recent studies the incidence of BM was shown to be significantly lower in PCI than in respective control arms. NVALT11/DLCRG-02 study addressed the occurrence of symptomatic BM as an endpoint. This was defined as BM identified during control MRI and the patient manifesting at least one of the following symptoms: vomiting, cognitive or affective disturbances, seizures, and focal neurologic symptoms. The rationale behind this was to show how PCI can be effective in preventing symptomatic BM in this specific cohort and thus preventing QoL deterioration (1). In the NVALT11/DLCRG-02 study 7% (PCI group) vs. 24.2% (controls) patients developed symptomatic BM yielding an absolute reduction of nearly 20% probability over time (1). In the RTOG 0214 there was no mention about whether BM being symptomatic or not, however, the 10-year BM rates equaled 16.7% in the PCI arm and 28.3% in the observation arm making it highly likely that the proportion of symptomatic BM was reduced in a proportional manner (6). Li et al. did not specify any subgroup of BM with symptoms either, but again, the 5-year brain recurrence rate was 15.6% in the PCI arm vs. 45.3% in no-PCI arm (29). Should this proportion be maintained for symptomatic BM reduction, PCI may have a marked impact on preventing the QoL decline due to BM-associated symptoms.

Patient status (WHO) or previous treatment (chemotherapy, surgery) had no impact in the NVALT11/DLCRG-02 study on the occurrence of BM in planned subgroup analysis (1). A different study performed by Sun et al., also in a planned subgroup analysis, reported that younger patients (<60 years old) and those with non-squamous disease develop BM more often than older patients and squamous histology (6). In the work by Li et al. such analyses were not performed or reported (29).

In both NVALT11/DLCRG-02 and RTOG 0214 trials, all patients had undergone surgery, radiotherapy and chemotherapy which were a standard of care at time of patients’ accrual (1,6). In the trial by Li et al. it was stated that no postoperative radiation therapy was performed in patients in whom N2 status was revealed in the pathology report (27).

None of those three studies reported molecular data such as the mutation status of the treated tumors. Similarly, there are no information about subsequent treatment of BM in the control arms of any of the abovementioned recent studies. All of those trials, as well as previous ones suffered from very low accrual.

Despite the substantial reduction of the risk of developing BM, and data suggesting a DFS prolongation as well, PCI is not free of its own risks. The major limiting factor is that PCI may lead to radiation-induced toxicity. Therefore, complication of PCI and BM may both exert a similarly negative impact on cognitive function. The specific types of neurocognitive deterioration associated with BM occurrence depend on their location and damage incurred to nearby regions of the brain. PCI on the other hand induces a specific type of deterioration, which manifests its impact typically 3 months after the procedure in the form of mild deterioration of cognitive functions.

Toxicity assessed by a physician is more and more often taken into consideration as an important secondary endpoint or being a point of interest of separate publications (30,31). Typically, neurologic but also non-neurologic symptoms are assessed in patients who underwent PCI. Neurologic and non-neurologic events which may develop after PCI and associated toxicity include typically mild cognitive decline and mild headaches. However, the fear of a very serious or life-threatening neurologic events is very prevalent and treatment-limiting. It is critical to note that the tests used to assess neurologic functions are not very specific and are one typically used for dementia screening. This is associated with limitations arising from the lack of validation of studies in other cohorts of patients with cognitive impairment. The most commonly used tools are mini mental state examination (MMSE) (32), Hopkins Verbal Learning Test (HVLT) (33) or Activities of Daily Living (ADL) (34).

Non-neurologic toxicity of PCI predominantly manifests as vomiting, which intensity dwindles with time, while the long-lasting neurologic toxicities exacerbate as time from treatment elapses. The non-neurologic sequelae of PCI are mainly assessed using the EORTC QLQ-C30 and brain-specific EORTC questionnaires (QLQ-BN20).

Evolution of adverse events overtime was a subject of the NVALT11/DLCRG-02 study. In the PCI arm of NVALT11/DLCRG-02 neurologic adverse events tended to develop overtime but were mild (grade 1 to 2). The worsening of QoL scores for the patients occurred during the time of treatment or soon after the treatment, mainly due to the appearance of immediate non-neurologic symptoms (1).

In the RTOG 0214 study over a 10-year follow up in terms of neurocognitive function the authors claimed that
there were insufficient data to conclude anything about long term neurocognitive toxicity (6).

An earlier pooled analysis of the RTOG 0214 data was performed on the basis of neurological evaluation with MMSE and HVL T. The results did not reveal any signs of a serious mental decline 1 year after PCI. Pooled analysis of RTOG 0214 and RTOG 0212 revealed that the deterioration of QoL may be considered as a good surrogate for poor OS (18).

Li et al. reported similar results: the main acute toxicities of a grade of at least 3, associated with PCI occurred within 90 days of treatment start. The reported complications included headache (in 1% of patients) and fatigue (2%). Severe headaches after 90 days were noted in 25% of the patients, fatigue in 1% and severe skin atrophy was observed in 1 individual. No significant differences between the two arms were shown in terms of QoL questionnaires.

The risk of neurologic symptoms (especially neurocognitive function decline) caused by PCI, may be potentially reduced by hippocampal avoidance (HA) (35) or the use of memantine (36). Preventing mental deterioration by using whole brain radiotherapy (WBRT)-as a standard technique used for PCI - plus memantine with or without HA showed, in a phase III trial, that patients with HA present no increased risk of worse control and HA protects from neurocognitive deterioration (36). Memantine plus HA was associated with lower risk of deterioration in contrast to control arm (HR =0.74, P=0.02) (36).

The NVALT11/DLCRG-02 study emphasized also patient-reported occurrence of adverse events. The numbers reported by patients in terms of vomiting, dizziness, headache, hypersomnia, memory impairment were in fact higher in PCI arm than in the observational arm (55 of 88 patients vs. 36 of 88 patients, correspondingly) but the grades of such outcomes was assessed as mild (1).

Given that the tests used for evaluation of neurologic deterioration performed in recent studies involving PCI are mostly adapted screening tests for cognitive impairment of any kind. This creates a major need for the wider use of other, more sensitive tools that should be used to assess mild neurologic deterioration or an effect of prophylaxis using memantine or HA in PCI. Such tests do exist but are still not validated in cohorts of patients with cancer. One such tool, designed for the assessment of mild memory and cognitive function impairment is a 10-minute cognitive screening tool—the Montreal Cognitive Assessment (MoCA) (37).

This test is designed to be used as a high-sensitivity tool that would identify impairment mild enough to slip through the MMSE test, with an intent for the MoCA to be used by primary care physicians. MoCA was already validated not only in US, Brazilian, and Asian populations with dementia but also on patients with stroke and in terms of neurological and cognitive rehabilitation effects. Another promising questionnaire is the Brain Health Assessment (BHA). This validated 10-minute, tablet-based test assesses aspects of neural performance other than memory (executive functions, speed, visuospatial skills or language), as these areas may deteriorate before overt memory failure develops (38). Given the improvements in survival rates and the pressure to preserve QoL in cancer survivors it becomes more and more reasonable to perform baseline cognitive performance assessment before initiating treatment (with or without PCI) to better monitor the patients. The abovementioned tools, or close collaboration with clinical psychologists or psychiatrists, will become more and more important in safeguarding patients from cognitive and neurological complications.

### Cost-effectiveness of preventing BM (PCI and other forms of preventing BM)

BM in patients diagnosed with NSCLC imposes a very high economic burden. It is therefore of utmost importance to analyse and compare different methods of BM prevention or treatment not only in terms of efficacy but also in terms of healthcare costs and their derivates. These may include outpatient and inpatient costs together with pharmacy costs, work absenteeism and potential dependence on social care. Those measures should be standardized (i.e., by average monthly expenditure per patient) and ideally should be adjusted for inflation in specific countries. Those kinds of analyses bring also a true light in real cost load of BM prevention and treatment. Cost-effectiveness analyses typically rely on cost utility analyses, where quality adjusted life years (QUALY) are primary outcomes. Those analyses should be standardized by appropriate reporting (39). According to American Society of Clinical Oncology (ASCO), there is currently highlighting on interventions demonstrating value instead of just effectiveness, which is of utmost importance for health care payer.

Thus far, analyses of cost reduction were performed mostly for molecularly targeted therapies and usually are performed on datasets obtained during the execution of clinical trials. This was the case for the analysis performed on the cohort assembled during the ALEX study (40,41).
The cost associated with BM treatment in abovementioned study was estimated as $6,029 per month over a 24-month period of total treatment. When the alectinib and crizotinib-treated groups were compared, it was shown that alectinib reduced those costs by approximately $41,000 per patient simply by reducing the incidence of BM. However, the analysis did not include the costs of additional alectinib treatment, and did not consider the additional expenditure associated with using alectinib instead of crizotinib (40).

So far, the cost-effectiveness of analysis or estimation of costs of PCI was not assessed in patients with NSCLC. De Ruysscher et al. claimed to evaluate the health costs but their analysis focused on the estimation of costs as associated with adverse events outcomes, as these are an integral component of such analyses, but no cost-effectiveness analyses were published from this cohort thus far (1). Cost-effectiveness analysis of PCI was performed in SCLC patients by means of a Markov model with internal validation, where PCI with HA (Arc therapy) was used. This treatment modality seemed to be more effective than conventional PCI assuming the risk of developing BM on level of 14% or by preventing neurocognitive dysfunction by at least 40% rates reduction (42). The two-way sensitivity analysis performed in this study revealed that PCI with HA would be the preferred strategy if every 10% decrease of neurocognitive dysfunction brings less expenses than increased risk of new metastases to 3% (42).

**Conclusions**

PCI may be of use as a tool of preventing BM, especially symptomatic ones, but it demands a very careful analysis of pros and cons with the patient. The impact of PCI on OS in NSCLC, even in highly selective groups of patients or with the use of neuroprotective techniques or supportive care will be marginal if any at all. However, the impact may be highly beneficial in terms of improved QoL if toxicity is avoided. Given the need for an individualized approach to PCI use, there is an urgent need to identify and define the group that has the greatest potential benefit from any kind of preventing BM. Without such guidelines it will be hard to optimize the use of PCI in NSCLC in an evidence-driven way. The effect of PCI in specific subgroups of patients: younger individuals with adenocarcinomas, staged III disease, patients with a high volume of disease, individuals with tumors with ALK translocations or EGFR mutations is particularly interesting as these individuals may benefit from group-specific effects of PCI and other preventative measures against BM. The use of protection or sparing brain from toxicity is also of interest—mainly through the use of memantine and HA (or at least administering radiation dose as low as reasonably acceptable in this organ at risk) by means of new techniques of radiotherapy. By minimizing toxicity one can expect a reduction of BM risk, with a potential benefit in terms of QoL and prolongation of DFS, or perhaps even OS in long term observations. Therefore, optimized and tailored reduction of BM risk seems to be a highly cost-effective treatment. Unfortunately, performing studies aimed at the optimization of PCI and toxicity prevention might be very difficult due to limited recruitment performance. Therefore, the most urgent need is to devise optimal strategies for recruiting individuals to BM-prevention trials and until high quality data are obtained, individualize the treatment approach when deciding upon PCI use in patients with cancer.

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**Future development of PCI—current ongoing studies**

There are several recently started, ongoing trials registered on clinicaltrials.gov:

(I) Prophylactic Cranial irradiation in erlotinib/gefitinib-responders with NSCLC—RT1001-trial identifier NCT01158170 (6)—this phase III trial, taking place in China, includes patients with NSCLC stage III after response to erlotinib/gefitinib treatment. The primary endpoint is BM incidence, secondary endpoints are OS, safety profile and tolerability, QoL. Status-unknown.

(II) Hippocampal-sparing prophylactic cranial irradiation in pathologically nodal positive non-small cell lung cancer—NCT02448992 (43). This phase II/III study will assess PCI with hippocampal spring vs. no PCI in patients with NSCLC stage III N+ after completion of chemotherapy. This study has already stopped recruitment. Its estimated completion date is December 2020. The primary endpoint is BM incidence, irrespective of symptoms, secondary endpoints are the effects of hippocampal-sparing PCI on neurocognitive functions (NCF).
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**Footnote**

*Conflicts of Interest:* The authors have no conflicts of interest to declare.

*Ethical Statement:* The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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