A phase Ib/IIa trial of 9 repurposed drugs combined with temozolomide for the treatment of recurrent glioblastoma: CUSP9v3


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Abstract

Background. The dismal prognosis of glioblastoma (GBM) may be related to the ability of GBM cells to develop mechanisms of treatment resistance. We designed a protocol called Coordinated Undermining of Survival Paths combining 9 repurposed non-oncological drugs with metronomic temozolomide—version 3—(CUSP9v3) to address this issue. The aim of this phase Ib/IIa trial was to assess the safety of CUSP9v3.

Methods. Ten adults with histologically confirmed GBM and recurrent or progressive disease were included. Treatment consisted of aprepitant, auranofin, celecoxib, captopril, disulfiram, itraconazole, minocycline, ritonavir, and sertraline added to metronomic low-dose temozolomide. Treatment was continued until toxicity or progression. Primary endpoint was dose-limiting toxicity defined as either any unmanageable grade 3–4 toxicity or inability to receive at least 7 of the 10 drugs at ≥ 50% of the per-protocol doses at the end of the second treatment cycle.

Results. One patient was not evaluable for the primary endpoint (safety). All 9 evaluable patients met the primary endpoint. Ritonavir, temozolomide, captopril, and itraconazole were the drugs most frequently requiring dose modification or pausing. The most common adverse events were nausea, headache, fatigue, diarrhea, and ataxia. Progression-free survival at 12 months was 50%.

Conclusions. CUSP9v3 can be safely administered in patients with recurrent GBM under careful monitoring. A randomized phase II trial is in preparation to assess the efficacy of the CUSP9v3 regimen in GBM.
Key Points

- Glioblastoma escapes pharmacological treatment as a result of cellular heterogeneity and resistance mechanisms.
- A treatment regimen with 9 different drugs (CUSP9v3) in addition to low-dose metronomic temozolomide was devised to tackle this issue.
- CUSP9v3 is safe in patients with recurrent GBM.

Importance of the Study

In 2013, we proposed a new concept to treat patients with recurrent GBM called Coordinated Undermining of Survival Paths (CUSP). The CUSP concept attempts to block growth-driving signaling pathways active in GBM. We took advantage of repurposing already-marketed non-oncological drugs and looked at the evidence for their ability to inhibit one or more of the identified GBM growth and cell survival pathways. Including pharmacology, drug interaction, and safety considerations, a list of 9 drugs was proposed to be used with low-dose, continuous temozolomide (CUSP9v3). Here, we report the results from the first clinical trial of CUSP9v3. In 10 patients with recurrent glioblastoma, the regimen was well-tolerated. This work is the first step in establishing that an extensive multi-drug regimen is tolerable and should now be tested for its potential efficacy against GBM.

As of fall 2020, current standard treatment of glioblastoma (GBM) with neurologically safe maximal resection, irradiation and temozolomide leads to progression-free survival (PFS) of 6.7 months, overall survival (OS) of 16.0 months, and 2-year OS of 30.7%. Recurrence usually takes place within a year after initial treatment. There is no commonly accepted standard of care for recurrent GBM. No regimen has proven to be safe and markedly effective for this condition.

In an attempt to address this unmet need, our group, together with many others, embarked on an exhaustive systematic search for already-marketed non-oncological drugs that might be able to set the stage for temozolomide to be more effective.

A complex winnowing process led to the final selection of the 9 drugs of Coordinated Undermining of Survival Paths combining 9 repurposed non-oncological drugs with metronomic temozolomide—version 3 (CUSP9v3) on which we report here the first clinical experiences. Details of that selection process can be found in the background papers.

Important criteria for drug selection were 1) robustness of preclinical data on GBM growth inhibition, 2) low side effect burden, 3) clinical familiarity with the drug in its general medicine (non-oncology) role, 4) availability as a generic, non-proprietary drug, and finally 5) lack of predictable serious pharmacological interactions.

The drugs of CUSP9v3 with their basic pharmacological attributes are listed in Table 1. Briefly, aprepitant inhibits NK-1 which is a growth-stimulating element in GBM. The anti-rheumatoid arthritis drug auranofin inhibits thioredoxin reductase, resulting in increased intracellular reactive oxygen species. The anti-hypertensive captopril reduces invasion, migration and adhesion of GBM cell activity through soluble matrix metalloproteinase (MMP)-2 and MMP-9 inhibition. The analgesic celecoxib has long been shown to have anticancer properties related to cyclooxygenase-2 inhibition and has demonstrated encouraging results in combination with low-dose temozolomide. The alcohol deterrent disulfiram is consistently cytotoxic to a wide range of cancer cells and is effective against GBM stem cells through aldehyde dehydrogenase inhibition. The antifungal itraconazole likely exerts its anticancer activity due to its multiple pharmacological effects with specific data in GBM pointing towards an effect on autophagy. The antibiotic minocycline has well-characterized neuroprotective effects and reduces GBM growth and invasion. The anti-retroviral ritonavir is effective in mouse GBM models with temozolomide by inducing endoplasmic reticulum stress. Last, the antidepressant sertraline was included for its ability to inhibit P-glycoprotein at the blood-brain barrier and because of its safe use in GBM patients. Overall, these drugs were judged to have robust anti-glioma or temozolomide-augmenting effects as well as to meet the criteria 1 to 5 above.

CUSP9v3 also comprises low-dose, continuous temozolomide at 20 mg/m² body surface area (BSA) p.o. twice daily, without interruption. This choice was based on past trials of various temozolomide schedules. After evaluating the 15 trials reviewed by Chen et al. and comparing these to data of Omuro et al., Clarke et al., and Reynés et al. who used temozolomide at 50 mg/m² BSA/day without interruption, that of Stockhammer et al. who used 20 mg/m² BSA/day and that of Zustovich et al. who used 40 mg/m² BSA/day, we concluded that any potential advantage of higher dosing was small and offset by a strongly reduced side effect burden associated with a regimen of 50 mg/m² BSA/day or less. Kong et al. reported that temozolomide at the dose of 40 mg/m² BSA/day was well-tolerated even in patients with Karnofsky Performance Status (KPS) < 70%. We, therefore, chose 20 mg/m² BSA given twice daily, the dose used by Zustovich et al. and Kong et al.
Although the safety profile of each drug of the CUSP9v3 protocol is well-known, safety concerns may arise due to the risk of drug-drug interactions at the pharmacodynamic (e.g., in form of additive toxicity) or pharmacokinetic level (with effects on metabolism or elimination, requiring dose adjustments or drug pausing). A database search prior to study initiation showed that clinically relevant interactions between CUSP9v3 drugs are expected to occur mainly due to CYP3A inhibition by itraconazole (strong), ritonavir (strong), and aprepitant (moderate). The unusual risks of using 10 daily drugs over a protracted period were partially offset by the good safety profile of each when used as a single drug and the intensity of our monitoring of patients.

We report here the results of the first trial of the CUSP9v3 regimen for recurrent or progressive GBM.

### Patients and Methods

#### Study Design

This is a phase Ib/IIa trial examining the safety of the CUSP9v3 regimen combined with temozolomide in patients with recurrent or progressive GBM. The primary endpoint was dose-limiting toxicity (DLT), and secondary endpoints were best tumor response, PFS, and OS. Dose-limiting toxicity was defined as either any unmanageable grade 3–4 toxicity at the end of the second treatment cycle or inability to receive at least 7 of the 10 drugs, all of them being given at ≥ 50% of the target doses, at the end of the second treatment cycle. Best tumor response was defined as the best therapeutic effect recorded from the start of the treatment until the last follow-up according to Response Assessment in Neuro-Oncology (RANO) criteria.29

Overall survival was defined as the time in months between the CUSP9v3 induction cycle start date and the date of last follow-up or death of any cause, whichever came first. Patients alive at the time of the last follow-up were censored.

Progression-free survival was defined as the time between the CUSP9v3 induction cycle start date and the date of the last follow-up, progression according to RANO criteria, or death of any cause, whichever came first. Patients with no progression and alive at the time of the last follow-up were censored.

This study was approved by the institutional review board of Ulm University Hospital (approval number 1 12/16) and the German competent authority Bundesinstitut für Arzneimittel und Medizinprodukte (BfArM; reference number 4041326) and registered at clinicaltrials.gov (NCT02770378).

### Table 1. Drugs Included in CUSP9v3 with Selected Pharmacological and Biological Characteristics

<table>
<thead>
<tr>
<th>Drug</th>
<th>p450 inhibition</th>
<th>Half-life</th>
<th>Core survival pathway or process targeted</th>
<th>Most frequent side effects according to drug label (in descending order of frequency)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aprepitant</td>
<td>3A4, 2C9</td>
<td>10 h</td>
<td>NK-1 receptors</td>
<td>Constipation, dyspepsia, fatigue, ALAT increase, decreased appetite, headache, hiccups</td>
</tr>
<tr>
<td>Auranofin</td>
<td>None</td>
<td>10 d</td>
<td>Thioredoxin, ROS generation, STAT3</td>
<td>Diarrhea, pruritus, exanthema</td>
</tr>
<tr>
<td>Captopril</td>
<td>None</td>
<td>2 h</td>
<td>ACE, AT1 receptors, MMPs</td>
<td>Diarrhea, nausea, dry mouth, constipation, abdominal pain, vomiting, loss of taste, dizziness, dysgeusia, sleep disorders, dyspnea, cough, rash, alopecia, pruritus</td>
</tr>
<tr>
<td>Celecoxib</td>
<td>2C9, 3A4</td>
<td>12 h</td>
<td>COX-1 and -2, carbonic anhydrase -2 and -9</td>
<td>Arterial hypertension</td>
</tr>
<tr>
<td>Disulfiram</td>
<td>2E1</td>
<td>&lt; 2 h</td>
<td>ALDH, ROS generation</td>
<td>Nausea, vomiting, drowsiness/somnolence</td>
</tr>
<tr>
<td>Itraconazole</td>
<td>3A4</td>
<td>19 h</td>
<td>P-gp efflux transporters, BCRP, hedgehog, 5-lipoxygenase</td>
<td>Nausea, abdominal pain, headache</td>
</tr>
<tr>
<td>Minocycline</td>
<td>None</td>
<td>10–20 h</td>
<td>Inhibits monocyte, macrophage and microglial contributions to growth</td>
<td>Diarrhea, nausea, vomiting, dyspepsia, flatulence, dizziness, rash, urticaria, pruritus</td>
</tr>
<tr>
<td>Ritonavir</td>
<td>3A4</td>
<td>4 h</td>
<td>P-gp efflux transporters (weak), proteasome, Akt, mTOR, cyclin D3</td>
<td>Pancreatitis, diarrhea, nausea, abdominal pain, vomiting, dyspepsia, fatigue, athenia, flushing, feeling hot, increased amylase, decreased thyroxine, arthralgia, back pain, dizziness, peripheral neuropathy, headache, paresthesia, dysgeusia, oropharyngeal pain, cough, pharyngitis, rash, pruritus</td>
</tr>
<tr>
<td>Sertraline</td>
<td>Weak</td>
<td>1 d</td>
<td>Akt, mTOR, TCTP</td>
<td>Diarrhea, nausea, dry mouth, fatigue, dizziness, drowsiness/somnolence, headache, insomnia, ejaculation failure</td>
</tr>
</tbody>
</table>

ACE, angiotensin-converting enzyme; Akt, protein kinase B; ALAT, Alanine aminotransferase; ALDH, aldehyde dehydrogenase; AT1, angiotensin II receptor type 1; BCRP, breast cancer resistance protein; COX, cyclo-oxygenase; MMPs, matrix metalloproteinases; mTOR, mammalian target of rapamycin; NK-1, neurokinin-1; P-gp, P-glycoprotein; ROS, reactive oxygen species; STAT3, signal transducer and activator of transcription 3; TCTP, translationally controlled tumor protein.
Sample Size

A sample size of 10 patients was selected to assess the primary endpoint. In this population, we expected a true rate of DLT of 40%. Sequential boundaries were used to monitor the DLT rate with accrual to be halted if excessive numbers of DLTs were seen. A Pocock-type stopping boundary yielded a probability of crossing the boundaries of maximally 10% when the actual rate of DLT was equal to the expected rate of 40%. The boundaries are described in Supplementary Material 1.

Patients

Eligible patients were adults with histologically confirmed GBM and recurrent or progressive disease according to RANO criteria. In 3 cases, study inclusion was allowed based on early recurrence that had not met minimal RANO requirements (ie, 10 mm x 10 mm diameters) but was judged as recurrence by external radiologists and confirmed by the trial's neuroradiologist (Be.S.). Patients with prior low-grade glioma were eligible if the malignant transformation to GBM was histologically confirmed. Additional key eligibility criteria were: no more than 3 prior episodes of tumor progression, KPS of at least 70%, stable steroid dose for at least 1 week prior to the start of study treatment, sufficient interval since last treatment (at least 4 weeks for systemic treatment or surgery, at least 12 weeks for radiotherapy) and no known contraindication to any of the CUSP9v3 drugs.

Treatment Regimen

Treatment initiation encompassed the addition of the 9 drugs to uninterrupted temozolomide as depicted in Supplementary Material 2 and comprised an induction cycle with 2 phases: a low-dose drug-by-drug addition phase followed by an up-dosing phase. Patients were hospitalized during the drug-by-drug addition phase, which lasted 18 days, to monitor tolerability and drug-drug interactions.

In summary, as schematically depicted in Supplementary Material 2, the treatment started with temozolomide (20 mg/m² BSA b.i.d.) and apreinpat (80 mg q.d.) on day 1, followed by the addition of 1 drug every 2 days (day 3, day 5 etc.) at the low-dose level. The last drug (auranofin) was added on day 17. On day 19, the up-dosing phase started with the dose of only one drug being increased every 2 days. The doses of temozolomide and apreinpat remained unchanged, 7 drugs were up-dosed only once and 1 drug (ritonavir) was up-dosed twice.

After reaching target doses of all drugs, the regimen remained unchanged until side effects mandated dose modifications and/or drug pausing or until tumor progression occurred. While the study was locked after the last recruited patient had completed 12 months of treatment, patients without tumor progression continued to receive the CUSP9v3 regimen beyond that point.

Safety and Dose Modifications

In addition to the potential drug-drug interactions to be monitored, we assessed the cumulative toxicity of the regimen. Employing the summary of product characteristics of each of the 10 drugs, we were able to identify the side effects most likely to occur during treatment. By developing a simple algorithm based on the frequency of each side effect (from very common [occurs in ≥ 1/10 patients] to very rare [occurs in < 1/10 000 patients]), we elaborated a strategy for dose modifications, dose re-escalations, and on-hold rules. For instance, in case of fatigue, no action would be taken for grade 1 or grade 2 fatigue. If grade 3 fatigue occurred, the first drug on a hierarchical list specific for fatigue was to be held until grade 2 or lower was reached and resumed at the same level. Additional drugs were to be held in the absence of resolution of symptoms.

During the induction cycle and the first 2 treatment cycles, adjustments (dose reductions and drug pausing) were allowed to accommodate the patients’ individual tolerability of the regimen. These modifications were discussed by a team comprising a neurosurgeon (M.-E.H.), an oncologist (R.M.-S.), a pharmacologist (O.Z.), and a psychiatrist (R.E.K.). For each patient, the regimen tolerated at the time of completion of the second treatment cycle (around day 90) was used to assess the primary endpoint.

Response Assessment

Response to study treatment was determined by neurological examination and contrast-enhanced magnetic resonance imaging (MRI) using the RANO criteria. Assessment was done at week 6, week 10 and then every 8 weeks. We used the best overall response, i.e., the best response recorded from the start of the treatment, as a secondary endpoint.

Statistical Analysis

Study data were analyzed by means of descriptive methods using frequencies (absolute and relative values) for categorical data as well as median and range for metric data. The Kaplan-Meier method was used to calculate PFS and OS. The median PFS and OS, respectively, are presented along with their corresponding 95% confidence intervals (CI). All analyses were performed using SAS (version 9.4, www.sas.com) and R (version 3.5.2, www.r-project.org).

Results

Patients Characteristics

Ten patients were included between August 2016 and April 2018. A total of 12 patients were screened. One patient could not be included because of high serum transaminases and one because of acute deep vein thrombosis.
Demographic characteristics of the 10 included patients are presented in Table 2.

Safety

The defined Pocock-type safety boundaries for stopping the trial were not crossed at any time. Nine patients completed at least 2 treatment cycles. At the end of the second treatment cycle, no patient had experienced any unmanageable grade 3–4 toxicity, and all patients had received at least 7 of the 10 drugs, given at ≥ 50% of the target doses. The primary endpoint was therefore met. Most frequently paused were ritonavir (for ataxia and fatigue), temozolomide (for diarrhea, nausea and laboratory abnormalities), captopril (for diarrhea and nausea), and itraconazole (for diarrhea and laboratory abnormalities) while ritonavir (for gait disturbance) and captopril (for fatigue) were most frequently dose-reduced.

All patients experienced at least one adverse event (AE) of any grade (Table 3) and 7 patients (70%) experienced at least one grade 3–4 AE, including 2 with at least one grade 4 AE. Ten AEs occurred in 5 or more patients. These AEs are presented in Table 3 with their grades and the drug(s) to which the respective AE was most likely related. All grade 3–4 AEs are presented in Table 4, grouped by classes, and listed with the drug(s) to which the AE was attributed or deemed possibly related. For all central nervous system AEs, no direct relationship could be established between the suspected drug and the respective AE because of the underlying disease.

All but one of the AEs attributed to CUSP9v3 drugs ceased upon pre-specified, targeted dose reduction or drug pausing within a range of 0 to 4 weeks (median 2 weeks; median number of drug modifications necessary to revert an AE: 1.5). None of the AEs that had ceased upon dose reduction or drug pausing recurred after the suspected drugs were reinstated.

Efficacy

Best overall response was stable disease (SD) in 6 patients and progressive disease (PD) in 4 patients (Table 5). Median duration of response was 8 (range 1–11) months in responders at the time of data lock. For 5 patients, SD was ongoing at the time of reporting. Three patients developed no detectable tumor on MRI during study treatment but would not be assigned “complete response” according to RANO because their tumor was “non-measurable” on MRI (ie, had maximal diameters of < 10 mm × 10 mm) at study entry. However, early recurrence had been initially diagnosed by external radiologists and was confirmed by the trial’s neuroradiologist (Be.S.) on the basis of a > 25% increase of “non-measurable” disease according to RANO.

Progression-free survival and OS are presented in Figure 1 and Figure 2, respectively. Both PFS and OS at 12 months were 50% with large confidence intervals because of the small sample (95% CI, 27–93%). Table 3 shows each patient’s individual PFS together with their disease characteristics and treatment prior to study entry.

Discussion

In this phase Ib/IIa trial in recurrent or progressive GBM, we found that 9 carefully selected non-oncological repurposed drugs together with twice daily 20 mg/m² BSA
temozolomide was safe and generally well-tolerated if individual dose adjustments were performed. CUSP9v3 is well-enough tolerable to be started in outpatients and to be fully introduced over a shorter time period than the 35 days that we used. The most frequent AEs are not expected to cause management problems as they are well-known to physicians treating patients with recurrent GBM.

Other trials in pediatric and adult high-grade glioma had reported the safety of various multi-agent regimens combining chemotherapy with repurposed drugs, using a range of 4–7 agents. Here we show that it is possible to combine 9 repurposed drugs given a careful evaluation of potential drug-drug interactions and cumulative toxicity. Knowing that many non-oncological drugs target pathways relevant to GBM, precision oncology approaches could expand their armamentarium by evaluating non-cancer drugs and combining them with classical cancer drugs. Such was recently reported in a trial in diffuse intrinsic pontine glioma. As strategies targeting cell membrane marker-defined glioblastoma cells may be limited, CUSP9v3 is consciously intended as a biomarker-independent approach.

During the protocol development for this trial, a hierarchical drug list had been developed (based on AE information contained in each drug’s summary of product characteristics) that correlated AEs to ranked sequences of drugs to be halved in dose or paused until pre-specified lower Common Toxicity Criteria of Adverse Events grades were reached. If that was not the case within 3 days, the next drug on the hierarchical list was halved in dose or paused etc. This strategy proved successful in managing AEs.

Noteworthy, the 3 AEs most frequently observed in this trial were nausea, headache and fatigue. While a causal relationship between the CUSP9v3 drugs and these cardinal AEs cannot be excluded, these symptoms may also be caused by the underlying disease itself and/or its primary treatment, temozolomide.

While the trial was not designed to assess the efficacy of the CUSP9v3 regimen, we observed that 5 patients progressed quickly, dying within a range of 1.5–7 months. The 5 other patients did well on treatment, all 5 having a PFS of 12 or more months (range 12–29 months at the time of data lock). In recurrent GBM, single-agent trials have reported PFS at 6 months of up to 20–30%. The rate of patients being alive and progression-free at 6 months has been suggested as an appropriate surrogate endpoint for predicting OS.

However, the small number of patients prevents any interpretation about the efficacy of the CUSP9v3 regimen. Another limitation is that the trial did not require histopathological confirmation of recurrence prior to study entry. Therefore, despite radiological judgement and reasonable time periods between the completion of radio- and chemotherapy on the one hand and the beginning of the study on the other hand (12 and 4 weeks, respectively), patients with a favorable course could have had pseudo-progression upon starting CUSP9v3. This issue of pseudo-progression is inherent to non-randomized trials in the recurrent setting.

One hypothesis that is supported by the pronounced dichotomy of response to CUSP9v3 is that CUSP9v3 may be more effective in patients with slower proliferating tumors and/or lower tumor burden, suggesting that this regimen may have a role in a prophylactic maintenance setting after first-line treatment.

Drug repurposing represents a large source of therapeutic options in cancer. In GBM in particular, notably 76 repurposed drugs recently were reported as potentially useful. The selection of the 9 drugs to be included in CUSP9v3 was a long iterative process within a conceptual framework that considered the specific and relevant preclinical,..
pharmacological, and empirical features of each drug in addition to the 5 criteria listed in the Introduction. It should not be assumed that combining other repurposed non-cancer drugs will automatically yield similar results; other regimens may prove more or less toxic or more or less effective.

In the common aggressive cancers, and especially in GBM, phenotypic spatial and temporal heterogeneity, in both stem and non-stem cell subsets, is a dynamic process responding to treatment interventions and driven further by hypoxia.43–45 In addition, GBM may be considered a collection of mutually interacting, mutually supporting cellular subpopulations46 demanding the use of a multi-drug combination to achieve prolonged treatment response.

**Conclusions**

We report here the first clinical trial of the CUSP9v3 regimen in recurrent or progressive GBM. The treatment regimen was safe under clinical, laboratory and electrocardiogram monitoring. A multicenter randomized controlled phase II/III trial is in preparation to assess the efficacy of the CUSP9v3 regimen in GBM.
Table 5. Tumor Characteristics, Prior Treatment and Outcomes on CUSP9v3 for Each Patient

<table>
<thead>
<tr>
<th>Patient ID</th>
<th>Age at inclusion (years)</th>
<th>KPS at inclusion (%)</th>
<th>Prior treatment besides standard of care *</th>
<th>MGMT promoter status/IDH1/2 mutation</th>
<th>Best response</th>
<th>PFS (months)</th>
<th>Vital status at data lock</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>31</td>
<td>100</td>
<td>Re-resection</td>
<td>Methylated/mutated</td>
<td>SD</td>
<td>29</td>
<td>Alive</td>
</tr>
<tr>
<td>2</td>
<td>48</td>
<td>80</td>
<td>Re-resection, re-RT</td>
<td>Methylated/wild-type</td>
<td>PD</td>
<td>2</td>
<td>Deceased</td>
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<tr>
<td>3</td>
<td>60</td>
<td>70</td>
<td>Bevacizumab</td>
<td>Non-methylated/wild-type</td>
<td>PD</td>
<td>0</td>
<td>Deceased</td>
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<td>4</td>
<td>53</td>
<td>100</td>
<td>-</td>
<td>Methylated/wild-type</td>
<td>SD</td>
<td>21</td>
<td>Alive</td>
</tr>
<tr>
<td>5</td>
<td>41</td>
<td>100</td>
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<td>Methylated/wild-type</td>
<td>SD</td>
<td>21</td>
<td>Alive</td>
</tr>
<tr>
<td>6</td>
<td>41</td>
<td>70</td>
<td>-</td>
<td>Methylated/wild-type</td>
<td>PD</td>
<td>0</td>
<td>Deceased</td>
</tr>
<tr>
<td>7</td>
<td>30</td>
<td>90</td>
<td>-</td>
<td>Non-methylated/wild-type</td>
<td>SD</td>
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<td>8</td>
<td>47</td>
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<tr>
<td>9</td>
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<td>10</td>
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<td>100</td>
<td>TTFields™</td>
<td>Non-methylated/wild-type</td>
<td>SD</td>
<td>12</td>
<td>Alive</td>
</tr>
</tbody>
</table>

* All 10 patients had been treated with surgery, chemo-radiotherapy and adjuvant temozolomide.

ID, identification; IDH1/2, isocitratdehydrogenase 1 or 2 gene; KPS, Karnofsky Performance Score; MGMT, O6-methylguanine-DNA methyltransferase; PD, progressive disease; PFS, progression-free survival; RT, radiotherapy; SD, stable disease.

Figure 1. Progression-free survival since CUSP9v3 start.
This pilot study met its primary endpoint with no un-
predicted side effects resulting from the combination of
drugs, and there was a signal of a potential positive effect
of CUSP9v3 that should be tested in future trials.

Supplementary Material
Supplementary material is available at Neuro-Oncology
Advances online.

Keywords
chemotherapy | clinical trial | drug repurposing | glioblas-
toma | multi-drug combination

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Authorship Statement. M.E.H., R.E.K., G.B., and K.B. con-
A.D., T.H., O.Z., Be.S., A.S., L.M., L.B., Bi.S., M.D.S., and
M.-A.W. were involved in the conduct of the trial. B.M., T.H.,
M.E.H., and R.E.K. analyzed and interpreted the data. M.E.H.,
R.E.K., G.B., and T.H. wrote the manuscript. All authors re-
viewed the manuscript.

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