Abstract: The standard of care for treatment of glioblastoma results in a mean survival of only 12 to 15 months. Convection-enhanced delivery (CED) is an investigational therapy to treat glioblastoma that utilizes locoregional drug delivery via a small-caliber catheter placed into the brain parenchyma. Clinical trials have failed to reach their endpoints due to an inability of standard catheters to fully saturate the entire brain tumor and its margins. In this study, we examine the effects of controlled catheter movement on dye dispersal volume in agarose gel brain tissue phantoms. Four different catheter movement control protocols (stationary, continuous retraction, continuous insertion, and intermittent insertion) were applied for a single-port stepped catheter capable of intrainfusion movement. Infusions of indigo carmine dye into agarose gel brain tissue phantoms were conducted during the controlled catheter movement. The dispersal volume ($V_d$), forward dispersal volume ($V_{df}$), infusion radius, backflow distance, and forward flow distance were quantified for each catheter movement protocol using optical images recorded throughout the experiment. $V_d$ and $V_{df}$ for the retraction and intermittent insertion groups were significantly higher than the stationary group. The stationary group had a small but significantly larger infusion radius than either the retracting or the intermittent insertion groups. The stationary group had a greater backflow distance and lower forward flow distance than either the retraction or the intermittent insertion groups. Continuous retraction of catheters during CED treatments can result in larger $V_d$ than traditional stationary catheters, which may be useful for improving the outcomes of CED treatment of glioblastoma. However, catheter design will be crucial in preventing backflow of infusate up the needle tract, which could significantly alter both the $V_d$ and shape of the infusion.

Keywords: convection-enhanced delivery; dispersal volume; glioblastoma; catheter movement

1. Introduction

Glioblastoma is a highly infiltrative brain tumor with margins that typically extend centimeters beyond the visible tumor [1]. Even with treatment, patient survival is poor, with a five-year survival rate of 5.5% [2] and a median survival of 12 to 15 months [3]. Gold-standard treatment protocols involving surgical resection, radiotherapy, and concomitant or adjuvant chemotherapy are not effective and recurrence is unavoidable [4–6]. Even combinatory therapy, which uses multiple standard-of-care treatments, has resulted in only moderate improvement in overall patient survival [7], and the tumor generally recurs at or near the site of the original tumor location [8,9].

The presence of the blood–brain barrier (BBB) and blood–brain–tumor barrier (BBTB) make systemic delivery of therapeutics to the brain challenging. In order to bypass the BBB and overcome the limitations of diffusive delivery of drugs, Bobo et al. proposed the use of convection-enhanced delivery
CED uses locoregional delivery of therapeutics to bypass the BBB by inserting a small caliber catheter though a burr hole in the skull and directly into the brain parenchyma. This technique utilizes pressure-driven flow to deliver large amounts of therapeutics directly to the region of interest [10, 11]. CED infusions occur primarily within the interstitial space of the brain. Finally, drug infusion times can range from a few hours to several days [12].

Although treatment of malignant gliomas using CED appeared promising after phase I [13–15] and phase II [14] clinical trials, a phase III (PRECISE) trial failed to meet its clinical endpoints [16]. A retrospective study conducted by Sampson et al. examined possible reasons for the failures of the PRECISE (Phase III Randomized Evaluation of CED of IL13-PE3QQR with Survival Endpoint) trials and concluded that inadequate drug delivery volume to the tumor may have been an important indicator for the failure of the trial, as on average only 20.1% of the 2 cm penumbra surrounding the tumor resection cavity was covered by the drug [17]. Additionally, initial trials of CED treatment for Parkinson’s disease using a glial cell-line-derived neurotrophic factor (GDNF) showed promising results [18, 19]. However, larger clinical trials failed to reach their primary endpoints [20], and a retrospective study found that only 2–9% of the target area was covered [21].

Standard catheters are unable to fully penetrate target volumes with CED treatments and have additionally been shown to be susceptible to reflux up the needle track, ultimately reducing the overall infusion volume [22, 23]. In order to combat the low delivery volumes found in clinical trials, focus was directed to improving the catheter design. Many of these devices focused on limiting backflow using a step change [24–28] and adding multiple ports within a single primary cannula to increase total infusion volume [29, 30]. However, exploration of alternative methods to increase the volume dispersed ($V_d$), such as catheter movement during infusion, has been sparse. Breeze et al. initially proposed the use of retraction in order to achieve an elongated deposition while implanting fetal tissue in the putamen of patients with Parkinson’s disease [31]. No other groups have attempted to recreate the elongated geometry with continuous retraction but some clinical trials have relied on depositing multiple payloads of therapeutics to target the putamen to treat Parkinson’s disease [32–34]. Additionally, Sillay et al. examined two-step advancement and retraction of catheters in agarose phantoms [35]. The study indicated that advancement of the catheter resulted in less backflow and more spherical delivery than retraction. However, their results were dependent on the catheter geometry and they relied on manual catheter placement. Further, the primary measures were the radius of infusion and backflow of infusate rather than total $V_d$. Bankiewicz et al. used single-step advancing catheters after an initial deposit in order to elicit backflow to reach over 50% of the putamen with a single catheter [36]. Further, Sudhakar et al. used an “infuse-as-you-go” technique, where real-time MRI was used to guide catheter advancements of between 2 and 4 mm in order to cover the putamen of non-human primates. “Infuse-as-you-go” covered nearly 15% more of the putamen in about 38% of the time compared to bilateral infusions via transfrontal trajectories [37]. Finally, Lewis et al. found that an increase in step length of the recessed step catheter would result in controlled reflux and an increase in $V_d$ [38]. Unlike the previous work mentioned, in this study we examine the effects of continuous remote retraction and insertion, intermittent insertion, and stationary catheter infusions on $V_d$ in agarose gel brain tissue phantoms.

## 2. Materials and Methods

Two 8-cm-long catheters were manufactured using 250 µm ID/375 µm OD fused silica capillary tubing (TSP250350, General Separation Technologies, Inc., Wilmington, DE, USA). The capillary tubing was cleaved flat and attached to a 22 gauge plastic dispensing tip using cyanoacrylate glue and UV-cured acrylic. Further, using a similar process to work previously conducted [39, 40], a 50 µm solid core multimode optical fiber (M14L02, Thorlabs Inc., Newton, NJ, USA) was adhered to the annular wall of the capillary tube using cyanoacrylate glue, such that the two parts co-terminated. The addition of the multimode fiber made the catheters capable of both fluid and light energy delivery; however, the light delivery functionality was not examined in the present work.
Agarose gel has previously been shown to be a good model for healthy porcine brain tissue [41]; therefore, it was used as a brain tissue surrogate in this study. Agarose gel brain phantoms were prepared by creating an agarose solution (0.6% w/w) by mixing certified molecular biology agarose powder (Bio-Rad Laboratories, Hercules, CA, USA) and deionized water. The solution was heated to a low boil and continuously stirred until the mixture became optically clear. After a cooling period of 15 min, approximately 50 mL of agarose was poured into polystyrene molds, covered, and allowed to completely cool to room temperature (25 ± 2 °C).

Immediately prior to infusions, a custom acrylic lid with an integrated reflux arresting tube was placed on top of the agarose mold, such that the reflux arresting tip was inserted into the agarose, as shown in Figure 1. The reflux arresting tip was constructed from polyether-ether ketone (PEEK) tubing (1.01 mm ID/1.57 mm OD). This design allowed for the catheters to move relative to both the agarose and the reflux arresting tip, with both components remaining as fixed reference points.

![Diagram of shadowgraphy setup](image)

**Figure 1.** Schematic of shadowgraphy setup, where a linear positioner controls a stage used to move the catheter through a reflux arresting cannula into an agarose specimen. The manual stage allows for gross alignment of the catheter prior to insertion into the gel. A constant-flow syringe pump is used to deliver indigo carmine dye through the catheter and into the gel, while a camera is used to acquire images throughout the infusion.

A shadowgraphy setup similar to that used previously [29] allowed visualization of the dye $V_d$. Briefly, two agarose gel phantoms were placed on an acrylic stage in front of a light-diffusing shade and behind a complementary metal-oxide-semiconductor camera (Rebel T1i, Cannon Inc., Ohta-ku, Tokyo, Japan), allowing for two experiments to be conducted at the same time. The agarose gel phantoms were kept at 25 °C, consistent with previous studies [29,38,41] and below the gel’s gelling temperature of 36 °C. Two catheters were supported by two linear stages in series—the first stage allowed for manual deployment, while the second stage allowed for remote deployment, as shown in Figure 1. For all groups, the catheters were first manually inserted into the gel phantoms through the reflux arresting tubing using the manual linear stage, such that they just began protruding from the reflux arresting tip. Quick insertion rates were shown to minimize reflux in agarose gel phantoms [42]; therefore, once the catheters were safely guided into the phantoms, they were rapidly deployed at 600 mm/min to their starting point. The catheters were controlled with a high-precision linear
actuator (A-LAR-E series, Zaber Technologies, Vancouver, BC, Canada) via a Bowden cable attached to the linear stage containing the catheters. The actuator was controlled with a custom graphical user interface created using LabView (National Instruments, Austin, TX, USA). Four experimental groups were examined: (1) a standard stationary catheter acting as the control, (2) continuously retracting catheter, (3) continuously advancing catheter, and (4) intermittently advancing catheter. The stationary and continuously retracting catheters were inserted to a depth of 30 mm, whereas the continuously and intermittent advancing catheters were inserted to a depth of 6 mm. For all four groups, a 1 mm “decoring retraction” was used in an attempt to remove any agarose that may have become lodged in the catheters during insertion. The deployment distance, intrainfusion movement rates, and movement distances are summarized in Table 1. After the catheters were inserted, an indigo carmine dye solution (1% w/w) was infused at 1 µL/min for 100 min using a programmable syringe pump (Chemyx Fusion 100, Chemyx, Austin, TX, USA). The flowrate of 1 µL/min was chosen in order to minimize the potential of backflow and is the rage of flowrates range typically used in CED infusion studies [16,24–30,39,40,43]. Once the infusion was started, the continuously retracting or advancing catheters were moved 25 mm at a rate of 0.25 mm/min and the intermittent advancing catheters were inserted an additional 6 mm with 1 mm of retraction every 16 min, for a total additional insertion of 25 mm. The infusion line pressure was measured at 2 Hz using gage pressure sensors calibrated from 0 to 30 psig (0 to 1551 mmHg) and 0 to 1 psig (0 to 52 mmHg) (26PC series, Honeywell International Inc., Golden Valley, MN, USA). Pictures were taken at the start of the experiment and retaken every minute throughout the 100 min infusion.

**Table 1.** Experimental group parameters.

| Group                     | Initial Deployment Distance | Movement Direction | Movement Speed | Movement Distance | Number of Replicates |
|---------------------------|-----------------------------|--------------------|----------------|-------------------|----------------------|
| Stationary (Control)      | 30 mm                       | None               | 0 mm/min       | 0 mm              | 7                    |
| Continuously Retracting   | 30 mm                       | Backward           | 0.25 mm/min    | 25 mm             | 8                    |
| Continuous Insertion      | 6 mm                        | Forward            | 0.25 mm/min    | 25 mm             | 14                   |
| Intermittent Insertion    | 6 mm                        | Forward            | 600 mm/min     | 25 mm (5 mm every 16 min) | 6                    |

At the conclusion of the infusions, the images were post-processed using Matlab (Mathworks, Natick, MA, USA), as described previously [29,44]. First, the images were imported into Matlab and cropped to encapsulate only the surrounding area of infusion in the agarose gel. Images were then converted to binary using a threshold value of 0.8% of stock concentration. Concentration was determined based on a calibration conducted by diluting stock 1% (w/w) dye solution in deionized water and preparing agarose gel standards at 10 levels from 0% to 1.0% concentrations (n = 6/level). Pictures of the standards were taken and imported into Matlab. Each image was then cropped to a small area of the standard, then images were converted to grayscale and normalized to the 0% standard. The mean for each standard level was taken and a linear curve-fitting algorithm was used to create the model, which is shown in Figure 2.

Once each image was converted to binary, images were visualized and any artifacts that were present on the image were manually removed. Artifacts were defined as any geometry present on the binary image that did not contribute to infusion volume. The artifacts primarily consisted of small amounts of debris on the specimen container or presence of the catheter far outside of the infusion cloud. Next, the first image from each infusion was scanned to find the location of the catheter tip. Additionally, the angle of the catheter relative to the gel was determined. Then, the volume of介质 $V_d$ was calculated for two regions: the full dispersal region ($V_d$) and forward dispersal ($V_{df}$). For all groups except the continuous retraction group, $V_{df}$ was defined as any infusion volume extending from the starting position of the catheter. For the continuous retraction group, $V_{df}$ was defined as any infusion volume...
extending from the current location of the catheter tip toward the proximal end of the catheter at its present location. The location of the catheter tip was estimated based on the starting location of the catheter, the time elapsed, and the angle of catheter insertion. The radius, backflow distance, and forward flow distance were also calculated. The forward flow distance was defined as the axial length of \( V_{df} \) and the backflow distance was the axial length of the infusion that did not contribute to \( V_{df} \). Additionally, pressure and temperature readings were analyzed and experiments were discarded if either infusion line pressure eclipsed 50 mmHg or if the temperature strayed from a \( 25 \pm 2 \) °C window. During the experiment, one of the temperature sensors began malfunctioning (affecting 3 infusions). However, since two separate infusions were conducted adjacent to one another at the same time and under the same conditions, the temperature from the adjacent infusions were used to evaluate exclusion criteria when temperature malfunctions occurred. A one-way analysis of variance was conducted to test for significant \( V_{df} \) \( V_{dif} \) infusion radius, backflow distance, forward flow distance, and average pressure changes between groups. The analysis of variance was followed by a post-hoc Tukey’s test.

![Figure 2. Grayscale intensity values plotted against 1% (w/w) indigo carmine dye dilutions in agarose gel used to determine the intensity threshold for converting grayscale images into binary images.](image)

3. Results

A total of 35 experiments were conducted during this study. Seven stationary infusions were completed, with one outlier omitted from analysis because the pressure eclipsed 50 mmHg. Eight retraction infusions were performed, with two outliers omitted from analysis because the temperature strayed from the 2-degree window. Six intermittent insertion infusions were conducted, with none removed. Finally, 14 continuous insertion infusions were completed, with 13 omitted from analysis due to high infusion line pressure; due to lack of data (\( n = 1 \)), the entire group was removed from further analysis.

Volume Dispersed: A plot of \( V_d \) for each group after 100 min is shown in Figure 3a. The retracting group had the largest \( V_d \) (\( 1.33 \pm 0.05 \) cm\(^3\)) of any group and was significantly larger than both the intermittent insertion group (\( 1.09 \pm 0.14 \) cm\(^3\); \( p < 0.01 \)) and the stationary group (\( 0.88 \pm 0.11 \) cm\(^3\); \( p < 0.001 \)). The intermittent insertion group had the second largest \( V_d \) and also had a significantly larger volume than the stationary group (\( p < 0.05 \)).

Forward Volume Dispersed: The plot for \( V_{dif} \) is shown in Figure 3b. Both the retracting (\( 1.24 \pm 0.03 \) cm\(^3\); \( p < 0.001 \)) and intermittent insertion (\( 0.68 \pm 0.19 \) cm\(^3\); \( p = 0.001 \)) groups had significantly more \( V_{dif} \) than the stationary group (\( 0.34 \pm 0.13 \) cm\(^3\)), and the retracting group had significantly more \( V_{dif} \) (\( p < 0.001 \)) than the intermittent insertion group.

Backflow and Forward Flow Distance: The backflow and forward flow distances for each group are plotted in Figure 3c,d respectively. The retracting group (\( 0.71 \pm 0.16 \) cm\(^3\)) had less backward flow than the stationary group (\( 1.13 \pm 0.36 \) cm\(^3\); \( p < 0.05 \)). No statistical significance was found between the
intermittent insertion group (0.94 ± 0.09 cm$^3$) and the other two groups ($p > 0.2$). Finally, both the retracting (2.33 ± 0.03 cm$^3$) and the intermittent insertion (2.02 ± 0.08 cm$^3$) groups had larger forward flow distances than the stationary group (0.54 ± 0.11 cm$^3$; $p < 0.001$), with a larger forward flow distance in the retracting group than the intermittent insertion group ($p < 0.001$). In Figure 3d, the outlier was defined as 1.5 times the interquartile range under the first quartile. The outlier in the retraction group was calculated to be 1.6 times the interquartile range under the first quartile.

**Figure 3.** Plots of the (a) total volume dispersed ($V_d$); (b) forward volume dispersed ($V_{df}$); (c) backflow distance; (d) forward flow distance; and (e) infusion radius for the stationary, retraction, and intermittent insertion groups after 100 min of infusion. Note: * $p \leq 0.05$; ** $p \leq 0.01$; *** $p \leq 0.001$
Infusion Radius: The infusion radius for each group after 100 min is shown in Figure 3e. The stationary group (0.56 ± 0.03 cm) had a small but significantly larger infusion radius than either the retraction (0.48 ± 0.13 cm; p < 0.001) or the intermittent insertion (0.43 ± 0.03 cm; p < 0.001) groups.

Infusion Line Pressure: The infusion line pressure of the retraction group (9.62 ± 3.20 mmHg) was lower than the stationary group (14.43 ± 4.71 mmHg), but not significantly so (p < 0.08). The intermittent insertion group (12.49 ± 3.07 mmHg) showed no significant differences in infusion line pressure from the other groups.

Data over infusion time: Figure 4 shows the outcome measurements at 20-min intervals through the 100-min infusion for each group. The retraction group had a higher V_d than either the stationary or intermittent insertion groups for all calculated time points, with significant levels (p < 0.01) of V_d at all time points after 20 min. The intermittent insertion group had a significantly higher V_d than the stationary group at the 80 and 100 min marks only (p < 0.01). Additionally, the retraction group had significantly greater forward volumes than either of the other groups at all time points (p < 0.001), and the intermittent insertion group had significantly greater V_df than the stationary group only at the 80- and 100-min time points (p < 0.01). The stationary group had significantly higher infusion radii than the other groups at all time points greater than 60 min (p < 0.05), and the retraction group had significantly greater infusion radii than the intermittent insertion group at all time points after 20 min (p < 0.05). The stationary group only had significantly more backflow than the retraction group at the 100 min mark (p < 0.05). No significance was found for other groups or at other times. Lastly, the retraction catheter had significantly more forward flow distance than the other groups at all time points (p < 0.001). The stationary catheter had significantly more forward flow distance than the intermittent insertion group at the 20 min time point (p < 0.05), but the reverse is true for all other time points during the infusion (p < 0.001).
Figure 4. Plots of the (a) volume dispersed ($V_d$); (b) forward volume dispersed ($V_{df}$); (c) backflow distance; (d) forward flow distance; and (e) infusion radius for the stationary, retraction, and intermittent insertion groups at 20 min intervals throughout the 100 min infusion.

Continuous Insertion: In all of the continuous insertion experiments where catheter clogging occurred (13 of 14), significant tearing of the gel occurred, leaving fluid-filled voids in the agarose in all areas except the location of the catheter tip, as shown in Figure 5.

Figure 5. (a) Image of clogged continuously deploying catheter after 100 min of infusion. (b) Cross-sectional image of the same infusion after catheter removal.

4. Discussion

In this study, we examined the effect of stationary and moving catheters on the overall infusion volume, infusion radius, and backflow. We found a 1.5-fold increase in $V_d$ with the retracting group and a 1.2-fold increase with the intermittently inserted catheter compared to the stationary control.

Both of the catheter movement groups had significantly larger total $V_d$ values than the stationary group. The larger surface area over which infusion can occur created by catheter movement is likely the cause of the larger infusion volume; however, there is a tradeoff. The longitudinal infusion distance increases significantly with the moving groups, however the radius (horizontal infusion distance) decreases by a small but significant amount. Lewis et al. also found a significantly larger infusion volume and smaller infusion radii when using a recessed step catheter with large needle lengths placed before a reflux arresting step change, allowing for controlled reflux [38].

The reduction in radii found with both the retracting and intermittently inserted protocols could benefit from the use of catheters with multiple independent ports, such as the arborizing catheter that we previously developed [29]. A catheter with this design provides a reflux arresting geometry at the tip of the primary cannula and allows for multiple needles to be placed near each other. If the arborizing angle is created such that a small amount of overlap between individual needles occurs,
thus preventing voids in the infusion, the increased volume associated with retracting catheters could
be utilized.

The overall backflow distance for the retracting catheter was found to be significantly smaller
than the backflow distance of the stationary catheter. This finding is in contrast to results found by
Sillay et al. [35]. This could be due to the presence of the reflux-arresting feature used in this study.
For both the intermittent insertion and retracting groups, the reflux-arresting feature was located
approximately 5 mm from the catheter locations at the beginning and end of the infusions respectively.
The close proximity of the reflux-arresting feature could be responsible for reducing the measured
backflow distance. However, it should be noted that the backflow distance was statistically the same
for all groups at earlier time points. Another possible cause for this apparent discrepancy is the method
in which the backflow was calculated. Sillay et al. only reported backflow distances in groups where
the forward flow distance was smaller than the backflow distance, while we reported all backflow
distances. However, all of the stationary specimens had smaller forward flow distances than backflow
distances, and for all groups the backflow distances were larger than the radii, with the exception of
one retracting specimen.

In all specimens, dye never escaped the gel phantom. Further, dye was effectively arrested at or
before the reflux arresting step change in all but two intermittent insertion specimens. The amount of
reflux was also likely limited by the low flowrate chosen for the experiment, as reflux in the retracting
catheter group did not reach the reflux arrestor until late in the experiment. The low flowrate and
the quick insertion of the catheters at the beginning of the experiment likely limited agarose damage,
as we expect dye would quickly reflux up the entire needle tract as soon as catheter movement began.
While quick insertion rates limit damage to the agarose, and therefore reflux, Casanova et al. showed
that slow insertion rates in rat brains result in less tissue damage and less backflow [45,46]. If both
slow insertion speeds and slow retraction speeds are consistently used during continuous catheter
retraction treatments, tissue damage may be mitigated and backflow can be reduced.

The infusion pressure for the retracting group was noticeably, but not significantly, lower than
both the stationary and the intermittent insertion groups. High flowrate infusions have been shown to
result in increased backflow [22,23]. However, the decreased pressure associated with the retracting
catheters may allow for larger flowrates to be permissible, without the creation of backflow. An increase
in the flowrate without overcoming a reflux-arresting step could have a considerable impact on the
clinical relevance of large-volume CED infusions.

All but one of the continuous insertion experiments resulted in clogging of the catheter. Interestingly,
other studies have started flow during the insertion process in order to limit clogging [27,35,36,47,48].
However, the insertion rate used in this experiment was much lower than that of other experiments,
which may be partially responsible for these findings. Agarose buildup at the tip of the catheter was
consistent with the slow insertion speeds in a study conducted by Casanova et al. [42]. Casanova et al.
进一步 found that the build-up of agarose (agarose damage) reduced the prestress on the tissue,
allowing for more backflow. Given this finding and the presence of the large cavity in the agarose, it is
likely that most of the infusion volume in this group would have been backflow, even if the catheter
did not consistently clog.

Study Limitations: While agarose gel tissue phantoms have been used extensively to conduct
CED experiments and have been shown to act as surrogates to the interstitial space in the brain
tissue due to the gel’s poroelastic nature [49,50] and similar pore size to that of the brain [51],
agarose gel tissue phantoms do not model either the cells or the blood vessels that infusions will
typically follow [49]. Agarose gel tissue phantoms are homogenous and isotropic, which cannot
capture the infusion morphology of pressure-driven flow inside of the heterogeneous and anisotropic
brain. Additionally, the specific interactions of the catheter with the agarose gel during insertion or
movement may not mirror that of the brain [48]. This hypothesis has been proven to a certain extent,
as Casanova et al. demonstrated an inverse trend of tissue damage based on insertion speed in agarose
gel compared to brain tissue [42,45]. However, it should be noted that studies utilizing multistep
collinear payloads [32,37] did not note any adverse events due to intraoperative catheter movement. Nonetheless, additional work should be conducted that assess both the safety and efficacy of controlled catheter movement in vivo.

5. Conclusions

In this study, we assessed the effect of catheter movement on $V_d$ in agarose gel phantoms. Catheters were inserted into phantoms and either kept stationary, continuously retracted, continuously deployed, or intermittently deployed during the infusion. Images of the infusion were taken and processed in order to determine the $V_d$ of the infusion. Significantly higher $V_d$ was found with both the retraction (1.5-fold) and the intermittent insertion (1.2-fold) groups compared to stationary catheters. The use of dynamic catheters could not only improve tumor coverage but could also allow for the creation of patient specific infusion geometries, furthering the efficacy of CED treatment.

Author Contributions: Conceptualization, J.N.M. and C.G.R.; data curation, J.N.M.; formal analysis, J.N.M.; funding acquisition, C.G.R.; investigation, J.N.M. and G.R.M.; methodology, J.N.M. and C.G.R.; project administration, C.G.R.; resources, C.G.R.; software, J.N.M.; validation, J.N.M. and G.R.M.; writing—original draft, J.N.M.; writing—review and editing, J.N.M., G.R.M., and C.G.R. All authors have read and agree to the published version of the manuscript.

Funding: This research was funded by the National Institutes of Health and the National Cancer Institute, grant number P01 CA207206-01.

Acknowledgments: The authors would like to thank Vanessa Garlepp and Egleide Y. Elenes for their help in specimen preparation and data analysis.

Conflicts of Interest: The fabrication methods and applications of the catheter are described in U.S. Patent 10,220,124, issued on 5 March 2019, and managed by the Virginia Tech Intellectual Properties (VTIP) Group.

References

1. DeAngelis, L.M. Brain Tumors. *N. Engl. J. Med.* 2001, 344, 114–123. [CrossRef] [PubMed]
2. Ostrom, Q.T.; Gittleman, H.; Xu, J.; Kromer, C.; Wolinsky, Y.; Kruchko, C.; Barnholtz-Sloan, J.S. CBTRUS Statistical Report: Primary Brain and Other Central Nervous System Tumors Diagnosed in the United States in 2009–2013. *Neuro Oncol.* 2016, 18, v1–v75. [CrossRef] [PubMed]
3. Wen, P.Y.; Kesary, S. Malignant Gliomas in Adults. *N. Engl. J. Med.* 2008, 359, 492–507. [CrossRef] [PubMed]
4. Berger, M.S. Malignant astrocytomas: Surgical aspects. *Semin. Oncol.* 1994, 21, 172–185. [PubMed]
5. Paulsson, A.K.; Holmes, J.A.; Peiffer, A.M.; Miller, L.D.; Liu, W.; Xu, J.; Hinson, W.H.; Lesser, G.J.; Laxton, A.W.; Tatter, S.B.; et al. Comparison of clinical outcomes and genomic characteristics of single focus and multifocal glioblastoma. *J. Neurooncol.* 2014, 119, 429–435. [CrossRef] [PubMed]
6. Shaw, E.J. Central Nervous System Overview. In *Clinical Radiation Oncology*; Gunderson, L.L., Tepper, J.E., Eds.; Churchill-Livingstone: Philadelphia, PA, USA, 2000; pp. 314–354.
7. Stupp, R.; Mason, W.P.; van den Bent, M.J.; Weller, M.; Fisher, B.; Taphoorn, M.J.; Belanger, K.; Brandes, A.A.; Marosi, C.; Bogdahn, U.; et al. Radiotherapy plus concomitant and adjuvant temozolomide for glioblastoma. *N. Engl. J. Med.* 2005, 352, 987–996. [CrossRef]
8. Wallner, K.E.; Galicich, J.H.; Krol, G.; Arbit, E.; Malkin, M.G. Patterns of failure following treatment of glioblastoma multiforme and anaplastic astrocytoma. *Int. J. Radiat. Oncol. Biol. Phys.* 1989, 16, 1405–1409. [CrossRef]
9. Hochberg, F.H.; Pruitt, A. Assumptions in the radiotherapy of glioblastoma. *Neurology* 1980, 30, 907–911. [CrossRef]
10. Bobo, R.H.; Laske, D.W.; Akbasak, A.; Morrison, P.F.; Dedrick, R.L.; Oldfield, E.H. Convection-enhanced delivery of macromolecules in the brain. *Proc. Natl. Acad. Sci. USA* 1994, 91, 2076–2080. [CrossRef]
11. Morrison, P.F.; Laske, D.W.; Bobo, H.; Oldfield, E.H.; Dedrick, R.L. High-flow microinfusion: Tissue penetration and pharmacodynamics. *Am. J. Physiol.* 1994, 266, R292–R305. [CrossRef]
12. Vandergrift, W.A.; Patel, S.J.; Nicholas, J.S.; Varma, A.K. Convection-enhanced delivery of immunotoxins and radioisotopes for treatment of malignant gliomas. *Neurosurg. Focus.* 2006, 20, E13. [CrossRef] [PubMed]
13. Kunwar, S.; Prados, M.D.; Chang, S.M.; Berger, M.S.; Lang, F.F.; Piepmeier, J.M.; Sampson, J.H.; Ram, Z.; Gutin, P.H.; Gibbons, R.D.; et al. Direct intracerebral delivery of cintredekin besudotox (IL13-PE38QQR) in recurrent malignant glioma: A report by the Cintredekin Besudotox Intraparenchymal Study Group. J. Clin. Oncol. 2007, 25, 837–844. [CrossRef] [PubMed]

14. Patel, S.J.; Shapiro, W.R.; Laske, D.W.; Jensen, R.L.; Asher, A.L.; Wessels, B.W.; Carpenter, S.P.; Shan, J.S. Safety and Feasibility of Convection-enhanced Delivery of Cotara for the Treatment of Malignant Glioma: Initial Experience in 51 Patients. Neurosurgery 2005, 56, 1243–1253. [CrossRef] [PubMed]

15. Sampson, J.H.; Akabani, G.; Archer, G.E.; Bigner, D.D.; Berger, M.S.; Friedman, A.H.; Friedman, H.S.; Herndon Li, J.E.; Kunwar, S.; Marcus, S.; et al. Progress report of a Phase I study of the intracerebral microinfusion of a recombinant chimeric protein composed of transforming growth factor (TGF)-α and a mutated form of the Pseudomonas exotoxin termed PE-38 (TP-38) for the treatment of malignant brain tumors. J. Neuro Oncol. 2003, 65, 27–35. [CrossRef]

16. Kunwar, S.; Chang, S.; Westphal, M.; Vogelbaum, M.; Sampson, J.; Barnett, G.; Shaffrey, M.; Ram, Z.; Piepmeier, J.; Prados, M.; et al. Phase III randomized trial of CED of IL13-PE38QQR vs. Gliadel wafers for recurrent glioblastoma. Neuro Oncol. 2010, 12, 871–881. [CrossRef]

17. Sampson, J.H.; Archer, G.; Pedain, C.; Wembacher-Schroder, E.; Westphal, M.; Kunwar, S.; Vogelbaum, M.A.; Coan, A.; Herndon, J.E.; Raghavan, R.; et al. Poor drug distribution as a possible explanation for the results of the PRECISE trial. J. Neurosurg. 2010, 113, 301–309. [CrossRef]

18. Gill, S.S.; Patel, N.K.; Hotton, G.R.; O’Sullivan, K.; McCarver, R.; Bunnage, M.; Brooks, D.J.; Svendesen, C.N.; Heywood, P. Direct brain infusion of gial cell line-derived neurotrophic factor in Parkinson disease. Nat. Med. 2003, 9, 589–595. [CrossRef]

19. Slevin, J.T.; Gerhardt, G.A.; Smith, C.D.; Gash, D.M.; Kryscio, R.; Young, B. Improvement of bilateral motor functions in patients with Parkinson disease through the uniteral intraputaminal infusion of gial cell line-derived neurotrophic factor. J. Neurosurg. 2005, 102, 216–222. [CrossRef]

20. Lang, A.E.; Gill, S.; Patel, N.K.; Lozano, A.; Nutt, J.G.; Penn, R.; Brooks, D.J.; Hutton, G.; Moro, E.; Heywood, P. Randomized controlled trial of intraputamenal gial cell line–derived neurotrophic factor infusion in Parkinson disease. Ann. Neurol. 2006, 59, 459–466. [CrossRef]

21. Salvatore, M.F.; Ai, Y.; Fischer, B.; Zhang, A.M.; Grondin, R.C.; Zhang, Z.; Gerhardt, G.A.; Gash, D.M. Point source concentration of GDNF may explain failure of phase II clinical trial. Exp. Neurol. 2006, 202, 497–505. [CrossRef]

22. Chen, M.Y.; Lonser, R.R.; Morrison, P.F.; Govermale, L.S.; Oldfield, E.H. Variables affecting convection-enhanced delivery to the striatum: A systematic examination of rate of infusion, cannula size, infusate concentration, and tissue–cannula sealing time. J. Neurosurg. 1999, 90, 315–320. [CrossRef] [PubMed]

23. Morrison, P.F.; Chen, M.Y.; Chadwick, R.S.; Lonser, R.R.; Oldfield, E.H. Focal delivery during direct infusion to brain: Role of flow rate, catheter diameter, and tissue mechanics. Am. J. Physiol. 1999, 277, R1218–R1219. [CrossRef] [PubMed]

24. Krauze, M.T.; Saito, R.; Noble, C.; Tamas, M.; Bringas, J.; Park, J.W.; Berger, M.S.; Bankiewicz, K. Reflux-free cannula for convection-enhanced high-speed delivery of therapeutic agents. J. Neurosurg. 2005, 103, 923–929. [CrossRef] [PubMed]

25. Vazquez, L.C.; Hagel, E.; Willenberg, B.J.; Dai, W.; Casanova, F.; Batich, C.D.; Sarntinoranont, M. Polymer-coated cannulas for the reduction of backflow during intraparenchymal infusions. J. Mater. Sci. Mater. Med. 2012, 23, 2037–2046. [CrossRef] [PubMed]

26. Yin, D.; Forsayeth, J.; Bankiewicz, K.S. Optimized cannula design and placement for convection-enhanced delivery in rat striatum. J. Neurosci. Methods 2010, 187, 46–51. [CrossRef]

27. Gill, T.; Barua, N.U.; Woolley, M.; Bienemann, A.S.; Johnson, D.E.; Sullivan, S.O.; Murray, G.; Fennelly, C.; Lewis, O.; Irving, C.; et al. In vitro and in vivo testing of a novel recessed-step catheter for reflux-free convection-enhanced drug delivery to the brain. J. Neurosci. Methods 2013, 219, 1–9. [CrossRef]

28. Rosenbluth, K.H.; Luz, M.; Mohr, E.; Mittermeyer, S.; Bringas, J.; Bankiewicz, K.S. Design of an in-dwelling cannula for convection-enhanced delivery. J. Neurosci. Methods 2011, 196, 118–123. [CrossRef]

29. Elenes, E.Y.; Rylander, C.G. Maximizing Local Access to Therapeutic Deliveries in Glioblastoma. Part II: Arborizing Catheter for Convection-Enhanced Delivery in Tissue Phantoms. In Glioblastoma; De Vleeschouwer, S., Ed.; Exon Publications: Brisbane, Australia, 2017. [CrossRef]
30. Vogelbaum, M.A.; Brewer, C.; Barnett, G.H.; Mohammadi, A.M.; Peereboom, D.M.; Ahluwalia, M.S.; Gao, S. First-in-human evaluation of the Cleveland Multiport Catheter for convection-enhanced delivery of topotecan in recurrent high-grade glioma: Results of pilot trial 1. J. Neurosurg. 2018, 1–10. [CrossRef]

31. Breeze, R.E.; Wells, T.H.; Freed, C.R. Implantation of fetal tissue for the management of Parkinson’s disease: A technical note. Neurosurgery 1995, 36, 1044–1048. [CrossRef]

32. Marks, W.J., Jr.; Ostrem, J.L.; Verhagen, L.; Starr, P.A.; Larson, P.S.; Bakay, R.A.; Taylor, R.; Cahn-Weiner, D.A.; Stoessl, A.J.; Olanow, C.W. Safety and tolerability of intraputaminal delivery of CERE-120 (adeno-associated virus serotype 2–neurturin) to patients with idiopathic Parkinson’s disease: An open-label, phase I trial. Lancet Neurol. 2008, 7, 400–408. [CrossRef]

33. Marks, W.J., Jr.; Bartus, R.T.; Siffert, J.; Davis, C.S.; Lozano, A.; Boulis, N.; Vitek, J.; Stacy, M.; Turner, D.; Verhagen, L. Gene delivery of AAV2-neurturin for Parkinson’s disease: A double-blind, randomised, controlled trial. Lancet Neurol. 2010, 9, 1164–1172. [CrossRef]

34. Bartus, R.T.; Herzog, C.D.; Chu, Y.; Wilson, A.; Brown, L.; Siffert, J.; Johnson, E.M., Jr.; Olanow, C.W.; Mufson, E.J.; Kordower, J.H. Bioactivity of AAV2-neurturin gene therapy (CERE-120): Differences between Parkinson’s disease and nonhuman primate brains. Mov. Disord. 2011, 26, 27–36. [CrossRef] [PubMed]

35. Sillay, K.; Hinchman, A.; Kumbier, L.; Schomberg, D.; Ross, C.; Kubota, K.; Brady, M.; Brodsky, E.; Miranpuri, G.; Raghavan, R. Strategies for the Delivery of Multiple Collinear Infusion Clouds in Convection-Enhanced Delivery in the Treatment of Parkinson’s Disease. Stereotact. Funct. Neurosurg. 2013, 91, 153–161. [CrossRef] [PubMed]

36. Bankiewicz, K.S.; Sudhakar, V.; Samaranch, L.; San Sebastian, W.; Bringas, J.; Forsayeth, J. AAV viral vector delivery to the brain by shape-conforming MR-guided infusions. J. Control. Release 2016, 240, 434–442. [CrossRef] [PubMed]

37. Sudhakar, V.; Naidoo, J.; Samaranch, L.; Bringas, J.R.; Lonser, R.R.; Fiandaca, M.S.; Bankiewicz, K.S. Infuse-as-you-go convective delivery to enhance coverage of elongated brain targets. J. Neurosurg. 2019, 1, 1–8. [CrossRef]

38. Lewis, O.; Woolley, M.; Johnson, D.E.; Fletcher, J.; Fenech, J.; Pietrzyk, M.W.; Barua, N.U.; Bienemann, A.S.; Singleton, W.; Evans, S.L. Maximising coverage of brain structures using controlled reflux, convection-enhanced delivery and the recessed step catheter. J. Neurosci. Methods 2018, 308, 337–345. [CrossRef]

39. Hood, R.L.; Ecker, T.; Andriani, R.; Robertson, J.; Rossmeisl, J.; Rylander, C.G. Augmenting convection-enhanced delivery through simultaneous co-delivery of fluids and laser energy with a fiberoptic microneedle device. In Proceedings of the SPIE BIOS, San Francisco, CA, USA, 20 March 2013; p. 85760G.

40. Hood, R.L.; Andriani, R.T.; Ecker, T.E.; Robertson, J.L.; Rylander, C.G. Characterizing Thermal Augmentation of Convection-Enhanced Drug Delivery with the Fiberoptic Microneedle Device. Engineering 2015, 1, 344–350. [CrossRef]

41. Chen, Z.J.; Gillies, G.T.; Broadus, W.C.; Prabhu, S.S.; Fillmore, H.; Mitchell, R.M.; Corwin, F.D.; Fatouros, P.P. A realistic brain tissue phantom for intraparenchymal infusion studies. J. Neurosurg. 2004, 101, 314–322. [CrossRef]

42. Casanova, F.; Carney, P.R.; Sarntinoranont, M. Influence of needle insertion speed on backflow for convection-enhanced delivery. J. Biomech. Eng. 2012, 134, 041006. [CrossRef]

43. Vogelbaum, M.A. Convection enhanced delivery for treating brain tumors and selected neurological disorders: Symposium review. J. Neurooncol. 2007, 83, 97–109. [CrossRef]

44. Sabliov, C.M.; Boldor, D.; Keener, K.M.; Farkas, B.E. Image Processing Method to Determine Surface Area and Volume of Axi-Symmetric Agricultural Products. Int. J. Food Prop. 2002, 5, 641–653. [CrossRef]

45. Casanova, F.; Carney, P.R.; Sarntinoranont, M. In vivo evaluation of needle force and friction stress during insertion at varying insertion speed into the brain. J. Neurosci. Methods 2014, 237, 79–89. [CrossRef] [PubMed]

46. Casanova, F.; Carney, P.R.; Sarntinoranont, M. Effect of needle insertion speed on tissue injury, stress, and backflow distribution for convection-enhanced delivery in the rat brain. PLoS ONE 2014, 9, e94919. [CrossRef] [PubMed]

47. Hood, R.L.; Andriani, R.T., Jr.; Emch, S.; Robertson, J.L.; Rylander, C.G.; Rossmeisl, J.H., Jr. Fiberoptic microneedle device facilitates volumetric infusate dispersion during convection-enhanced delivery in the brain. Lasers. Surg. Med. 2013, 45, 418–426. [CrossRef] [PubMed]
48. Sillay, K.; Schomberg, D.; Hinchman, A.; Kumbier, L.; Ross, C.; Kubota, K.; Brodsky, E.; Miranpuri, G. Benchmarking the ERG valve tip and MRI Interventions Smart Flow neurocatheter convection-enhanced delivery system’s performance in a gel model of the brain: Employing infusion protocols proposed for gene therapy for Parkinson’s disease. *J. Neural. Eng.* 2012, 9, 026009. [CrossRef]

49. Raghavan, R.; Mikaelian, S.; Brady, M.; Chen, Z.-J. Fluid infusions from catheters into elastic tissue: I. Azimuthally symmetric backflow in homogeneous media. *Phys. Med. Biol.* 2009, 55, 281. [CrossRef]

50. Chen, Z.J.; Broaddus, W.C.; Viswanathan, R.R.; Raghavan, R.; Gillies, G.T. Intraparenchymal Drug Delivery via Positive-Pressure Infusion: Experimental and Modeling Studies of Poroelasticity in Brain Phantom Gels. *IEEE Trans. Biomed. Eng.* 2002, 49, 85–96. [CrossRef]

51. Gillies, G.; Allison, S.; Tissue, B. Positive pressure infusion of fluorescent nanoparticles as a probe of the structure of brain phantom gelatins. *Nanotechnology* 2002, 13, 484. [CrossRef]