Revisiting intra-arterial drug delivery for treating brain diseases or is it “déjà-vu, all over again”? 

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Abstract

For over six decades intra-arterial (IA) drugs have been sporadically used for the treatment of lethal brain diseases. In recent years considerable advance has been made in the IA treatment of retinoblastomas, liver and locally invasive breast cancers, but relatively little progress has been made in the treatment of brain cancers. High resting blood flow and the presence of the blood-brain barrier (BBB), makes IA delivery to the brain tissue far more challenging, compared to other organs. The lack of advance in the field is also partly due to the inability to understand the complex pharmacokinetics of IA drugs as it is difficult to track drug concentrations in sub-second time frame by conventional chemical methods. The advances in optical imaging now provide unprecedented insights into the pharmacokinetics of IA drug and optical tracer delivery. Novel delivery methods, improved IA drug formulations, and optical pharmacokinetics, present us with untested paradigms in pharmacology that could lead to new therapeutic interventions for brain cancers and stroke. The object of this review is to bring into focus the current practice, problems, and the potential of IA drug delivery for treating brain diseases. A concerted effort is needed at basic sciences (pharmacology and drug imaging), and translational (drug delivery techniques and protocol development) levels by the interventional neuroradiology community to advance the field.

Key words: Brain diseases, drug delivery, intra-arterial

INTRODUCTION

For over six decades intra-arterial (IA) drugs have been proposed for the treatment of lethal brain diseases. Recent reports of super-selective intraarterial (IA) injections of bevacizumab for treating human gliomas have brought IA drug delivery back into focus. In contrast to treatment of brain cancers the interest in IA chemotherapy for treating liver and breast cancers has considerably increased in the last three decades. Major advances have also been made in using IA chemotherapy of retinoblastomas to avoid surgical removal and improve function. High resting cerebral blood flow and the presence of the blood brain barrier (BBB), makes IA drug delivery to the brain far more challenging compared to other organs.

In the last two decades endovascular surgery has opened new vistas for treating neurological diseases. With easy access to distal cerebral circulation provided by modern micro-catheters, and finite diffusion distances within the capillary networks, one would expect that it would be easy to demonstrate the effectiveness of IA drugs. Unfortunately, this is yet to be realised or worked out. The failure in treating brain diseases with IA drugs is in part due to a continuing lack of understanding of IA pharmacokinetics, unavailability of convenient high-speed drug concentration measurement methods, poor rationalisation of drug injection protocols and the use of drugs and formulations without adequate pharmacokinetic justifications –many of which were never developed for IA use in the first place.

Yet, in-extremis, IA drugs are used for the treatment many other brain diseases beyond cancers, such as stroke, cerebral vasospasm, raised intracranial pressure, etc.
and infections.\[26,27\] Several anaesthetic drugs have been safely been used in diagnostic neurology for localisation of brain functions, since 1949, which reflects the safety of IA delivery.\[28-33\] In recent years, monoclonal antibodies, stem cells, viral vectors, liposomes and nanoparticles have been delivered by IA injections - all of which could play a major role in the future treatment of brain diseases.\[34\] Improved IA delivery could enhance onset and effectiveness of smart nano-scale drug delivery vehicles that are in development. Considering the practice, pitfalls, and the potential of IA drugs, this review makes the case for reassessing IA drugs and it elaborates on the novel treatments that IA drugs could offer. A concerted effort is needed at basic sciences (pharmacology and drug imaging), and translational research (drug delivery techniques and protocol development) levels by the interventional neuroradiology community to advance the field.

THE CHECKERED HISTORY OF IA DRUGS

In 1929, motivated in part by the desire to deliver drugs, Werner Forssmann demonstrated on himself the feasibility of cardiac catheterisation.\[35\] Two decades later Wada introduced IA anaesthetic drugs for localising brain functions and the Wada test is still used as a diagnostic tool for epilepsy treatment.\[28\] In the 1950s, Klopp et al., pioneered the use of intracarotid nitrogen mustard to treat brain cancers.\[1,36,37\] In the 1960s, Charles Wilson systematically investigated chemotherapy of brain tumours including IA delivery.\[38-40\] In the 1970s, Stanley Rapoport introduced intracarotid osmotic agents for disrupting the blood-brain barrier (BBB).\[41\] In the 1980s, Oldfield’s group at the National Institutes of Health (NIH) made a concerted effort to improve intracarotid chemotherapy and went as far as using extracorporeal haemofiltration to minimise systemic toxicity.\[42-45\] Unfortunately, there was little clinical impact of even such intense interventions while others reported unexplained focal neurological complications during IA infusions.\[46-50\] Consequently, the interest in IA drugs waned by the early 1990s.\[51\] Edward Neuwelt was one of the few people who continued the work, using IA mannitol for BBB disruption.\[52\] In hindsight though, the failures in 1980s were particularly unfortunate as they happened at the dawn of the modern endovascular era.\[53,54\]

THE FUNDAMENTAL PRINCIPLES OF IA DRUG DELIVERY

In simulations of steady-state IA infusions, Dedrick et al., showed that IA drug delivery was most effective when there is: (i) low regional blood flow; (ii) high regional extraction; and (iii) high systemic clearance; and that the advantage of IA delivery wanes with prolonged infusions [Figure 2].\[55\] Other investigators at the time pointed to the problem of “streaming”. Streaming occurs with low volumes drug infusions, when the currents within the arterial stream determine the distribution of the drugs. Maldistribution due to streaming could result in toxic concentrations in some territories while bypassing others.\[56-59\] Streaming was probably responsible for some of the focal neurological
complications reported with IA chemotherapy and it also explains some of the failures of pharmacokinetic models of IA drug delivery. The Dedrick model also reveals the problems in IA drug delivery to the brain, due to the presence of BBB which limits drug uptake, and the high resting blood flow to the organ. Technological advances in endovascular surgery now provide new opportunities to improve IA drug delivery by transiently reducing blood flow and injecting bolus of drugs as predicted by the Dedrick model. Injecting drugs as boluses during transient cerebral hypoperfusion not only enhances arterial concentrations, it avoids streaming and helps to localise drug interventions at specific sites.

It is well accepted that under normothermic conditions the brain can withstand 3 minutes of ischaemia. Several investigators have used transient flow arrest to better localise drug delivery. In 2004, Yamane et al., demonstrated the safety and efficacy of balloon occlusion during treatment of 187 paediatric cases of retinoblastoma. No significant complications, except transient bradycardia, were reported in 563 treatments of balloon assisted super-selective ophthalmic artery infusions. Recently, Riina et al., used 3-min occlusion with 2-min reperfusion cycles to treat brain stem glioma with bevacizumab over a period of an hour with good tumour response at 24 hours. Arresting blood flow by transiently suppressing cardiac activity to localise cyanoacrylate glue has been used in the treatment of high flow cerebral arteriovenous malformations. In a series of cases Pile-Spellman and Young described the safe use of adenosine-induced cardiac arrest for embolisation of such malformations.

Therefore, faced with lethal brain diseases, it is not hard to justify the potential risk of stroke due transient cerebral hypoperfusion (TCH) or even flow arrest, if it can localise therapeutic agents and/or dramatically improve drug delivery. In experimental animals, reduction in blood flow can improve drug delivery three to ten fold compared to IA injections made during normal blood flow. During TCH, IA drug delivery can be further enhanced by optimising bolus injection parameters and by using drug formulations with rapid tissue uptake.

**Optimising bolus injections**

The objective of bolus design is to match the area under the arterial blood concentration-time curve to tissue uptake. During TCH, injecting drugs as boluses offers a number of advantages besides avoiding streaming. First, the bolus volume generates high arterial concentrations exposing the brain tissue to virtually pure drug. This decreases the contact with blood components and binding proteins to generate high free drug concentrations. Second, bolus injections during TCH increase the transit time through the cerebral circulation from 1 second to as long as 1-2 minutes in experimental animals. High concentrations of free drugs, the prolonged capillary contact and short inter-capillary diffusion distance, all combine to significantly improve drug delivery when TCH is combined with bolus injections. A good example of this is the cationic formulations of liposomes dramatic improvements (100 fold) in drug delivery are possible compared to intravenous delivery of the same liposomes.

In theory, one can manipulate three parameters of bolus injections: (i) Volume, (ii) concentration, and (iii) frequency. The volume of the bolus should be such that it ensures maximum capillary contact, i.e., able to displace blood in the arteries and the capillaries but not the veins. Excessive volume can result in rapid transit though the capillary networks to decrease the time available for drug uptake. The concentration of the drug in the bolus is determined by the brain tissue uptake. The frequency of bolus injection should ensure steady replenishment of drug in the capillary networks as the concentration of the drug declines due to tissue uptake. In addition, one has to consider the total dose of the drug, its local toxicity and whether repeat hypoperfusion bolus injection cycles will be necessary to treat the underlying pathology.

**Optimising IA drugs and formulations to improve first pass drug extraction**

It is often not realised that drugs are seldom screened or developed for IA administration. Most reports of IA treatments describe the off-label use of intravenous (IV) drugs. For maximum benefit from IA delivery, drugs have to be rapidly extracted during their first pass through the cerebral circulation. One of the reasons for the wide application of IA drugs for liver cancer
treatment is the high hepatic extraction. For example, hepatic extraction of floxuridine can be as high as 49 - 99% after hepatic artery infusion. However, drugs and drug formulations can also be optimised for first pass extraction by brain tissue. A good example of this, again are the liposomal formulations. Cationic magnetic liposomes are almost 15 times more effective with IA delivery compared to IV administration. When given intravenously, cationic liposomes are rapidly sequestered in non-target organs. However, when injected intraarterially during TCH, not only are the particles spared of sequestration by non-target organs but they also avoid exposure to plasma proteins which protects their surface charge. We have observed that IA injections of charge neutral liposomes have poor uptake by brain tissue while cationic liposomes are readily retained. Furthermore, bolus IA injections of cationic liposomes achieve three to four-fold-greater tissue concentrations during TCH compared to injections during normal blood flow.

Optimising drug design

The drug uptake across the BBB is related to the permeability surface area product (PSA) of a drug. By modifying the drug molecule, such as, decreasing polar groups or increasing aliphatic groups, one can increase the PSA. The approach was used to work brain selective drug delivery of chlorambucil tri-butyl ester after IV injection. A similar approach could augment the delivery of IA drugs.

Receptor and transporter conjugated drugs

Many drugs have been conjugated with transporter and receptor ligands to increase drug delivery across the BBB via these physiological mechanisms. For example, OX-26 monoclonal antibody to transferrin receptor (TFR-1) was developed to target drug delivery to the brain tissue after IV injections. The TFR-1 receptors trigger receptor-mediated endocytosis capable of delivering large drug cargos across the BBB. However, IV injections of these compounds were not very effective due to the non-targeted liver uptake. In the brain, the TFR-1 receptor is more abundantly expressed in tumours compared to the normal neuronal or glial tissue and could provide a valuable ligand for targeting chemotherapeutic drugs. Improved IA methods of drug delivery might be able to selectively target brain tumours.

Cationic carriers and IA drug delivery

Cationic peptides and drugs are capable of penetrating the BBB and their effective delivery could be further enhanced by TCH assisted delivery. Furthermore, nanoparticles have been already conjugated with low molecular weight protamine to enhance tissue uptake. Beside protamine there are other cationic carrier systems in the brain, such as for cationic albumin (sometimes called the super-carrier) that can also deliver drugs across the BBB. In recent years cationic cell penetrating peptides such as Trans-activator of transcription (TAT) derived from human immuno-deficiency virus have emerged as vehicles for drug delivery across the BBB. Dramatic improvements in IA drug delivery might be possible if drugs were conjugated to cationic carriers and injected during TCH in a manner similar to cationic liposomes. Preliminary data with TAT suggested that is indeed the case.

Novel IA drugs

Another approach to IA drug development could be the development of drugs that have exceedingly short biological half-lives, such as adenosine. In non-human primates in a head-to-head comparison between intracarotid sodium nitroprusside (T1/2 of 90s) and adenosine (T1/2 of 1s), IA sodium nitroprusside reduced systemic blood pressure but had no effect on cerebral blood flow (CBF) while adenosine in large doses did not affect the blood pressure but dramatically increased CBF. Adenosine provides a conceptual prototype of IA drugs, whose biological half-life is so short that it is significantly metabolised during transit through cerebral circulation to minimise systemic side effects. Such short-acting drugs will be invaluable when multiple drugs are used for dynamic IA interventions based on rapidly changing tissue needs.

Additional strategies for improving IA drug delivery

Besides balloon occluding catheters, CBF can also be reduced safely by induced hypothermia, deep IV anaesthesia, or by hyperventilation. Hypothermia being neuroprotective could be particularly useful to enhance safety and the duration of TCH. The method of choice for reducing blood flow, whether sustained or transient, will probably depend on the individual drug.

Systemic rescue

Oldfield et al., used extracorporeal haemofiltration of jugular venous blood to reduce systemic toxicity. Other investigators have used systemic antidotes concurrent with IA injections, such as thiosulphates, to neutralise re-circulating chemotherapeutic drugs. Classical forced alkaline or acid diuresis can also eliminate the recirculating drug. In recent years cyclodextrins have become available as non-specific chelating agents that could be used for the purpose reducing systemic toxicity.

Novel tools to investigate IA drugs

If we accept the proposition that there can be significant improvements to IA drug delivery then, we have...
to determine how best to rapidly advance the field. A fundamental hurdle to the understanding of IA drug delivery has been the inability to track drug concentration measurements in real-time in a typical laboratory or clinical setting. Chemical analysis of tissue samples does not provide site-specific histories. Although multiple tissue biopsies have been taken from the same animal to obtain time histories, there is cumulative injury to the tissue. Microdialysis is not readily feasible for IA studies if bolus injections are used due to the time required for dialysis equilibrium. Pharmacokinetic studies are feasible with magnetic resonance imaging and radioisotopes but logistics and cost can be prohibitive for high volume research.

In contrast, novel optical methods can help us better understand the pharmacokinetics of IA drugs. Optical methods, such as diffuse reflectance spectroscopy,[96-99] multi-spectral and hyperspectral imaging, spatial frequency domain imaging (SFDI),[100-102] confocal microscopy and spectroscopy, and even optical coherence tomography provide us with novel pharmacokinetic tools. Collectively, spatial and temporal resolution of optical methods is bound to generate novel insights in drug delivery and the mechanisms of drug resistance. It is generally held that future brain treatments will be based on macromolecules, such as antibodies, peptides, gene fragments, liposomes or nanoparticles. Such macromolecules can easily be tagged with optical tracers without altering their pharmacokinetic properties. For drugs with a stable spectrum, distinct from haemoglobin and deoxyhaemoglobin, ultra-fast tissue non-invasive drug concentrations measurements are feasible at several times per second with diffuse reflectance spectroscopy and multispectral imaging.[95] Diffuse reflectance spectroscopy can track multiple chromophores, that permits quantification of BBB permeability, using a tracer like indocyanine green, while tracking tissue drug concentrations.[103] Multispectral imaging could help to elucidate tissue surface drug concentrations for those that have wavelength specific excitation and emission at sampling rates of hundreds of times per second. Hyperspectral imaging could also provide a planar map of spectral changes after drug injection that could be used for determining tissue drug concentrations. Both confocal microscopy and hyperspectral imaging will require time for image acquisition; however, the former method may provide a pharmacokinetic tomogram post delivery while tissue spectroscopy may quantify free, bound and metabolised drug. In theory, in vivo optical pharmacokinetic tomography is possible in large tissue volumes (several cm) with SFDI but such application is still in development.[101,102] As nanoparticle-based drug delivery evolves “optical pharmacokinetic tomography” will be critical in mapping macromolecules whose diffusion is likely to be restricted in the highly compartmentalised brain tissue and not just by the BBB.[104]

**POTENTIAL IMPACT OF IA TREATMENTS**

Effective, timely and safe drug delivery to brain tissue is perhaps the most persistent and significant problem in translational neuroscience research. Most of the neuropharmaceuticals that are in development will not cross the BBB. William Pardridge therefore emphasised that drug discovery and drug delivery must go hand in hand.[105] The ability of IA drugs to target specific regions of the brain pathology, rapidly achieving therapeutic concentrations with minimal systemic exposure, with the possibility of administering multiple drugs at the same time, could open new avenues for the treatment for brain diseases. Due to the complex matrix of neurological injuries, future cures for brain diseases might require curative cocktails, not magic bullets. Administration of multiple drugs by the IV route is often limited because of drug interactions and additive side effects. However, multiple drugs could be administered as pulses via the IA route. Such an approach might offer innovative treatments. In the case of cerebral vasospasm a pulse of IA adenosine, a potent short acting vasodilator, might transiently dilate the spastic arteries to permit delivery of less potent but longer acting vasodilator, such as nicardipine.[106] In brain cancer treatment, IA drugs could offer chemo-ablative therapy for the core of the tumour that is devoid of any function and anti-mitotic anti-angiogenic therapy in the peripheral zone with salvageable functions. For stroke, multiple IA drugs might be injected while imaging injury parameters.[107] Such highly individualised and dynamic IA treatments could enable future molecular reconstructive neurosurgery that could improve the quality of survival not just provide a cure.

**CONCLUSIONS**

In conclusion, the advances in endovascular neurosurgery now permit the development of more sophisticated IA drug delivery protocols while those in optical engineering promise to reveal the hitherto ill-understood pharmacokinetics of IA drugs. It’s not “Déjà-vu, all over again!” as Mr. Yogi Berra said. To the contrary, novel delivery methods, improved IA drugs, and optical pharmacokinetics, present untested paradigms in pharmacology and offer hope to patients with intractable neurological diseases. A concerted effort is needed at basic sciences (pharmacology and drug imaging), and translational research (drug delivery techniques and protocol development) levels by the interventional neuroradiology community to advance the field.
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