Microencapsulated Dopamine (DA)-Induced Restitution of Function in 6-OHDA-Denervated Rat Striatum in vivo: Comparison Between Two Microsphere Excipients

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SUMMARY

Biodegradable controlled-release microsphere systems made with the biocompatible biodegradable polyester excipient poly [DL-lactide-co-glycolide] constitute an exciting new technology for drug delivery to the central nervous system (CNS). The present study describes functional observations indicating that implantation of dopamine (DA) microspheres encapsulated within two different polymer excipients into denervated-striatal tissue assures a prolonged release of the transmitter in vivo. Moreover, in this regard, the results show that there were clear cut temporal differences in the effect of the two DA microsphere formulations compared in this study, probably reflecting variations in the actual composition (i.e., lactide to glycolide ratio) of the two copolymer excipients examined. This technology has considerable potential for basic research with possible clinical application.

Key words: Microencapsulated dopamine; poly [DL-lactide-co-glycolide]; striatum; apomorphine; 6-hydroxydopamine.

INTRODUCTION

The main neurochemical characteristic of Parkinson's disease is a marked lesion of the nigro-striatal dopamine (DA) pathway. Current clinical management of Parkinson's disease relies strongly on substitution treatment with the DA precursor L-DOPA and/or direct DA agonists, administered via the parenteral route. In recent years several attempts have been made to reverse neurologically debilitating nigrostriatal DA deficiency by different alternative means of replacement therapy [1]. Although some advances have been reported with neuronal and adrenal medullary tissue transplants, this approach has not yet proven simple and effective to compensate for the underlying DA neuronal loss in humans with Parkinson's disease [9-11]. Intracerebroventricular infusion is another method currently being tested in experimental Parkinsonism [4] and in attempted replacement therapy of certain other neurodegenerative diseases [6]. Ideally, however, the delivery of substitution agents should be restricted specifically to the relevant damaged regions of the central nervous system (CNS).

Injectable drug delivery formulations - so-called microspheres - have recently been developed, in which drugs are microencapsulated within biocompatible and biodegradable copolymer excipients like poly [DL-lactide-co-glycolide] [3]. These co-polymers are from the same class of material which has been used for nearly two decades in resorbable surgical sutures. The copolymer microsphere formulation...
protects drugs from degradation and allows release of the drug from its excipient at a controlled rate for prolonged periods of time (for weeks, or even months) /3,20/. In view of its potential therapeutic usefulness, we have studied the microsphere methodology as a means of delivering DA to the rat central nervous system (CNS) in vivo. Specifically, our previous reports have indicated the feasibility of this approach and preliminary evidence suggested that intrastriatally implanted DA microspheres could, over long periods, counteract the postsynaptic DA receptor up-regulation resulting from unilateral 6-hydroxydopamine (6-OHDA) lesion of ascending dopaminergic neurons /13,14/. Thus, these initial results support the concept that implantation of DA microspheres could provide an advantageous method for achieving prolonged release of functionally significant amounts of DA into striatal tissue in vivo to substitute for subnormal levels of the endogenous transmitter.

Another interesting advantage of the microspheres is that the duration of drug release can be modified by manipulating the "shell" constituent copolymer biodegradation kinetics, e.g., by changing the ratio of lactide to glycolide in the formulation /20/. To this end, DA was microencapsulated in two copolymer excipients having different lactide to glycolide molar ratios. The object of this report is to compare these 2 different copolymer formulations. This is accomplished by comparing the duration of time that these intrastriatally implanted DA microspheres are able to reduce the extent of DA agonist-induced rotational behavior in unilaterally 6-OHDA-lesioned rats, used as a functional index of postsynaptic DA receptor denervation supersensitivity.

MATERIALS AND METHODS

Animals, surgery and behavioral testing

Twenty five male Sprague-Dawley rats (200-250 g; AB Laboratorietjänst, Sollentuna, Sweden) were unilaterally lesioned in the ascending median forebrain bundle (MFB) of monoamine neurons (coordinates A-P -4.3, L+1.4, D-V -8.7 from bregma, midline, and top of skull, respectively) /16/. The neurotoxin used for the purpose was 6-hydroxydopamine x HCl (6-OHDA; Sigma, St Louis, MO, USA), administered in a dose of 8 μg/4 μl saline vehicle (containing 0.1% ascorbate). Two weeks after lesion, the rats were challenged with the classical DA agonist apomorphine (0.1 mg/kg s.c.) and rotational responses were monitored in a computerized rotometer set-up /13/. Only rats in which the DA denervation has been successful (≥ 90% striatal DA depletion) will display strong contralateral rotation to apomorphine challenge in this model /8,21/. Rats responding to apomorphine with less than 400 contralateral rotations per 60 min within the first two weeks of testing were eliminated from the study (n = 7). Testing of the remaining 18 positive responders was then continued on a weekly basis until a stable level of rotation was established (max 10% difference between three consecutive sessions). This criterion was typically met about 6 weeks after the initial testing session (data not shown).

Polymers employed for encapsulation of DA

DA was encapsulated in two types of copolymer excipient. One copolymer had a 50:50 molar ratio of lactide to glycolide (referred to as DA 50:50). The other copolymer had a 65:35 molar ratio of lactide to glycolide (DA 65:35). It was predicted that, because of its higher lactide content, the 65:35 copolymer will take longer to biodegrade than the 50:50 copolymer, thus potentially affording a longer duration of delivery of DA in vivo.

Implantation of DA microspheres

Once rats reached a stable rotational baseline level to DA agonist challenge (cf. above) they were stereotaxically injected under light ether anesthesia with a suspension of DA microspheres (prepared immediately prior to injection). DA 50:50 microspheres (n = 5; 15 mg microspheres/50 μl saline) or DA 65:35 microspheres (n = 9; 30 mg microspheres/50μl) were implanted into two sites (3 μl-deposits, 2 levels/site) in the mediodentral striatum (A-P +0.7, L 2.3, D-V -4.5 and -5.5; A-P +0.2, L 2.3, D-V -4.5 and -5.5) /16/. DA 50:50 is predicted to
biodegrade twice as fast as DA 65:35 \cite{13}. To ensure that similar amounts of DA would be released per unit time the quantity of DA 65:35 microspheres was therefore twice that of the DA 50:50 microspheres. A separate group of control rats (n = 4) received corresponding injections of empty microspheres (50:50 polymer formulation; 30 mg microspheres/50 μl saline).

The injections were performed with a 10 microliter glass capillary tube (calibrated at 3 microliter intervals) connected to a standard Hamilton syringe (50 μl) via polyethylene tubing. The entire duration of injection was 3 min/site. Upon completion of the second infusion, the injection cannula was left in situ for an additional 60 seconds before being slowly retracted. The skin wound was closed with surgical clips and the animals allowed to recover from anesthesia (about 30-45 minutes thereafter). They were placed into the rotometers and their behavior recorded for the subsequent 2-3 hours. Starting one week after implantation of the DA microspheres, the rats were repeatedly tested for DA agonist-induced rotation on a weekly basis for 6-8 weeks.

**Histology**

At the termination of the 6-8 week schedule the rats were perfused with 5% glutaraldehyde and a series of consecutive sections (10 μm) of the striatum was incubated with anti-DA antiserum \cite{7,12}. The immunocytochemistry was performed with the standard avidin-biotin complex method, thus allowing visualization of tissue DA immunoreactivity.

**Statistics**

The data are presented as the total number of apomorphine-induced contralateral rotations summed over 60 minutes, means ± SEM. Statistical comparisons between post-implantation responses and pre-implantation baseline values (= obtained in the last apomorphine test session before implantation) were made employing Student's paired t-test. The results were expressed as the mean and standard error of the mean (SEM). Probability levels of 5% or less were considered statistically significant.

**RESULTS**

**Acute rotational response to DA microsphere implantation**

The 18 rats selected for microsphere implantation experiments had responded with the typical "two-peak" contralateral rotation pattern when initially challenged with apomorphine (0.1 mg/kg s.c.), a reliable indicator of > 90% DA lesion success \cite{8,21}. As previously reported \cite{14}, upon recovery from ether anesthesia (= 30-45 min after intrastratal implantation of DA microspheres) rats exhibited immediate contralateral rotational behavior with an amplitude comparable to that of apomorphine (0.1 mg/kg s.c.) but with longer duration. Rats receiving DA 65:35 microspheres displayed a somewhat more protracted response to the microsphere implantation but had a peak rotation amplitude similar to the DA 50:50 microsphere group \cite{14}. Animals implanted with empty microspheres did not display rotational behavior.

**Rotational responses to apomorphine in DA microsphere-implanted rats**

**DA 50:50 implants**

The total number of pre-implantation contralateral rotations for this group of rats was 553 ± 58 (n = 5). When compared to this value, the central striatal DA 50:50 microsphere implants resulted in a reduction to 397 ± 19, a 29% decrease in the total number of apomorphine-induced contralateral rotations 4 weeks post-implantation. During the subsequent two weeks the response to apomorphine challenge slowly approached the pre-microsphere baseline (Fig. 1). At 6 weeks there was a reduction to 497 ± 48, representing a non-significant 11% decrease in the total number of apomorphine-induced contralateral rotations.

**DA 65:35 implants**

The total number of pre-implantation contralateral rotations for this group of rats was 506 ± 26 (n = 9). The DA 65:35 microspheres resulted in a reduction to 319 ± 43, representing a 37%
decrease in the total number of apomorphine-induced contralateral rotations two weeks post-implantation. From then on and up to 8 weeks following implantation there was an average reduction to 375 ± 50, corresponding to a 26% decrease in the total number of apomorphine-induced contralateral rotations (Fig. 2). Since, as indicated above, the DA 65:35 polymer excipient was predicted to outlast the DA 50:50, it is interesting to note that the reduction in the number of apomorphine rotations from week 3 through week 8 remained rather stable with the former formulation whereas the apomorphine response in the DA 50:50 group was no longer significantly suppressed after week 5.

Sham implants

During the 6 week testing period post-implantation, rats that received empty microspheres did not display any significant change in the number of apomorphine-induced contralateral rotations when compared to the pre-implantation baseline (Fig. 3).

Immunocytochemistry

Micrographs of DA 50:50 are not presented in this report. However, previously we demonstrated that DA was present in DA 50:50 microspheres 3 weeks after implantation /14/. Our immunocytochemical observation of these microspheres 6 weeks after implantation revealed that the majority of the microspheres had biodegraded or were in the process of doing so. There were, however, sparse traces of DA immunoreactivity in some microspheres that had not fully biodegraded. In no case was the immunocytochemical reaction comparable to that of DA 65:35. In the DA 65:35 implant group, however, there were still microspheres containing respectable amounts of DA at 8 weeks (Fig. 4A-B). Examination of the micrographs in this latter group also reveals microspheres in the process of biodegrading at 8 weeks post-implantation (Fig. 4B). No DA immunoreactivity was present in the microspheres or in the striatal tissue of the "sham"
The present investigation demonstrates that implantation of microencapsulated DA in the central striatum can counteract, over substantial periods of time, apomorphine-induced rotational behavior in rats with chronic unilateral 6-OHDA lesions of ascending (nigrostriatal) dopaminergic neurons, thereby confirming our previous preliminary findings /14/. The attenuated responsiveness to DA agonist challenge in this experimental model is likely to be explained by a down-regulation of denervation-sensitized striatal postsynaptic DA receptors, following implantation of the in vivo DA-releasing microspheres /13,14/. Moreover, in this regard, the results show that there were clear cut temporal differences in the effect of the two DA microsphere formulations compared in this study, reflecting variations in the actual composition (i.e., lactide to glycolide ratio) of the two copolymer excipients examined.

The DA 50:50 and DA 65:35 excipients were predicted to biodegrade at different time intervals, while simultaneously unloading functionally significant amounts of the polymer-encapsulated DA. Our data support the validity of these predictions under in vivo conditions.
Fig. 3: The number of apomorphine-induced contralateral rotations in 6-OHDA-treated rats before and for 6 consecutive weeks following implantation of empty microspheres. Bars represent the mean ± SEM (n = 4). The pre-implantation baseline value (= obtained in the last apomorphine test session before implantation) was 532 ± 108. This value is shown at time 0. The number of apomorphine-induced contralateral rotations post implantation were as follows: 1 week 529 ± 97, 2 weeks 460 ± 73, 3 weeks 506 ± 66, 4 weeks 465 ± 92, 5 weeks 510 ± 86 and 6 weeks 536 ± 83. Abscissa, time in weeks; ordinate, accumulated contralateral rotations recorded for 60 minutes. There was no statistical difference in the number of contralateral rotations post-implantation compared to the pre-implantation baseline value.

Immediately after implantation into the central striatum, both the DA 50:50 and the DA 65:35 microsphere mixtures elicited strong contralateral rotation, considered to be mediated by postsynaptic DA receptor stimulation. However, this behavioral response vanished over a period of about 3-4 h, tentatively due to rapid receptor desensitization /13,14/. In the subsequent weekly test sessions the DA 50:50 microsphere group displayed a significantly reduced total number of apomorphine-induced contralateral rotations up to and including the 4th testing week post-implantation. However, in the DA 65:35 experiment the apomorphine response was reduced for at least 8 weeks. Even at this time the response to the DA agonist challenge was significantly lower than the pre-implantation baseline, suggesting that there was still enough DA remaining to maintain a reduction of the 6-OHDA-induced denervation supersensitivity. This interpretation is further supported by the immunocytochemical analysis, revealing the persistence of microspheres with intact or near-intact DA contents also at 8 weeks after implantation (cf. Fig. 4A-B). It is also interesting to note in this context that the time-course and extent of reduction in denervation supersensitivity produced by implantation of DA 65:35 microspheres is comparable with that reported after striatal grafting of adrenal medullary chromaffin tissue /19/.

Our results support the idea that the altered duration of in vivo release of encapsulated drug from resorbable polymer excipients is a function of the actual copolymer composition. There is convincing evidence from in vitro experiments that the rate of drug release from microspheres is very sensitive to variations in the excipient constituents /2,20/. Interestingly, this seems to apply also in vivo, the major difference being that in this case the time scale for microsphere biodegradation is now weeks/months rather than minutes/hours /20/. In the present study, the implanted 65:35 DA microspheres resulted in a
Fig. 4: Photomicrographs of frontal cryostat sections (10 μm) of the denervated striatum after implantation of DA 65:35 (A,B) or empty (C) microspheres. All the sections were incubated with anti-DA antiserum (1/500). The sections were processed for immunocytochemistry using the standard avidin-biotin-peroxidase complex method.

A) DA 65:35 microcapsules 8 weeks following implantation. Notice that the tissue on either side of the injection tract is intact. DA microspheres depicted by the immunocytochemical reaction are still within the striatum. x 160.

B) This photomicrograph is a higher magnification of A. A number of immunoreactive DA microspheres can be observed (arrows). There are some microcapsules in the process of degrading (arrow heads). x 640.

C) Empty microspheres (arrows) 6 weeks after implantation. Notice that these microspheres are totally devoid of immunoreactivity. x 640.
rather constant reduction in the apomorphine response from week 3 through week 8. Thus, it appears that even though the liberation of DA from these polymers relies on hydrolytic surface erosion in vivo, the rate of this process is dependent on the actual excipient composition so as to allow reasonable control of the release in the manufacturing step by, e.g., altering the lactide to glycolide ratio in the copolymer. This concept is also supported by the results with the 50:50 DA microspheres. Importantly, the present data equally support the biocompatibility of the microsphere formulations. It might be suggested that biodegradation through hydrolysis of the excipients used would be harmful to the brain tissue. If so, it could be expected that the striatal tissue would be damaged during the process and thus the rats would no longer be able to respond to apomorphine after termination of the biodegradation period. Our results clearly demonstrate that this is not the case. First, the immunocytochemical study revealed that introduction of the microspheres produced minimal (if any) tissue damage (cf. Fig. 4). Secondly, the DA 50:50 animals returned to pre-implantation baseline values, and thirdly, animals implanted with "sham" microspheres (without DA, but of the same shell composition) did not display any reduction in their apomorphine-induced rotational behavior over the entire 6-week period studied.

The present copolymer microsphere preparations appear to compare favorably with other reported polymeric excipient devices used in studies aiming for sustained release of DA. Other researchers have recently reported functional results similar to ours, using single, mm-thick ethylene-vinyl acetate (EV Ac) DA-containing copolymer matrix rods or discs /5,22/. As compared to the microsphere preparations, these latter polymer designs inevitably cause much more mechanical tissue damage by virtue of their size when implanted and removed /22/; indeed, aspiration of parts of the overlying neocortex was required in order to put the disc-shaped DA-entrapping polymer in position for release into the striatum /5/. Moreover, the EV Ac devices are non-resorbable and while, in one of the studies, only restricted gliosis was reported /5/, the chronic biocompatibility also remains to be established. Although the microspheres used in our studies cannot be removed once implanted, they cause minimal tissue damage and no additional manipulation of the experimental animal is necessary to get the DA within reach of the striatal tissue.

As referred to above, the DA 50:50 microsphere implants into the central portion of the striatum resulted in a clear-cut, ≥ 4-week long, blunting of the response to apomorphine challenge. In contrast, in a previous study, we found no attenuation in the rotational response to apomorphine 3 - 40 days after lateral striatal DA 50:50 microsphere implants /14/. The present results therefore represent another addition to the growing list of examples of functional heterogeneity of the striatum /15,17,18,23/.

Injectable microsphere formulations have earlier been demonstrated as a feasible means of delivering drugs at intended sites of action, for prolonged periods of time, at required rates and in proper therapeutic doses, to targets outside the blood brain barrier /20/. The present results indicate that DA microsphere preparations have the potential of being employed as a source of transmitter replacement also within brain tissue in vivo. These formulations allow sustained diffusion of the microencapsulated DA into the CNS at a controlled rate for predetermined periods of time, assure functional significance and at the same time appear to remain compatible with the host tissue. This new "slow-release", target-directed approach provides not only a useful tool in basic neuroscience but conceivably, after completing similar experiments in non-human primates, it may also be employed in the clinical management of neurodegenerative illness such as Parkinson's disease.

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