Chamber Cannulation in Nonhuman Primates

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PURPOSE. To determine the magnitude of ocular hypotony and the length of recovery time to 6 and 10 mm Hg IOP following anterior chamber (AC) cannulation.

METHODS. Bilateral IOP was recorded 500 times per second via telemetry immediately before, during, and immediately after AC cannulation with a 27-G needle in 10 different sessions at least 2 weeks apart in four male rhesus macaques (nonhuman primates; NHPs) aged 3- to 6-years old. Bilateral IOP was recorded continuously using a proven telemetry system while the NHPs were under general anesthesia during IOP transducer calibration experiments involving manometric control of IOP via AC cannulation, then continuously after the AC needles were removed until IOP recovered to precannulation levels. The change in IOP from baseline to AC cannulation was tested using the signed-rank test. The times necessary for IOP to recover to 6 and 10 mm Hg, respectively, were calculated.

RESULTS. Average precannulation IOP was 11.5 mm Hg and significantly decreased to an average of 2.3 mm Hg immediately following AC needle removal (P = 0.0156). On average, IOP recovered from 2.3 to 6 and 10 mm Hg in 32.4 and 63.7 minutes, respectively. Recovery times of IOP were not affected by repeated AC cannulations every 2 weeks.

CONCLUSIONS. Generally, IOP recovers relatively quickly after repeated AC cannulation, and did not result in extended duration hypotony. It is important to consider hypotony in animal experiments and clinical procedures involving AC cannulation and paracentesis when consideration of IOP or its effects is important.

Keywords: hypotony, anterior chamber cannulation, nonhuman primates, intraocular pressure

Anterior chamber (AC) cannulation is a common procedure in both research studies and clinical care. In research studies using animal models of ocular disease, AC cannulation is used for acute IOP control1–5 during imaging and other procedures, and paracentesis is used to collect aqueous humor samples.6 Anterior chamber paracentesis is a common clinical treatment to lower IOP acutely in acute angle closure and in central retinal arterial occlusion (CRAO).7–13

Frequent episodes of persistent hypotony following AC cannulations or paracentesis procedures could represent a confounding variable in animal studies investigating diseases such as glaucoma wherein IOP is a major risk factor. Hence, we investigated the magnitude of hypotony and time course of recovery to normal IOP levels following AC cannulation in nonhuman primates (NHP) with a continuous telemetry system and assessed changes in IOP recovery time after repeated cannulation.

In human subjects, ocular hypotony has been defined as IOP at or below 5 mm Hg,14,15 below 6.5 mm Hg,16 and even less than or equal to 8 mm Hg in previous studies.17 We defined hypotony for this study as a persistent IOP less than 6 mm Hg, as 6 mm Hg is within range of hypotony threshold used in previous studies, and it is lower than the IOP we typically measure in normal NHPs with telemetry. Most commonly, the causes of ocular hypotony are wound leak, retinal detachment, and insufficient aqueous humor production or an increase in aqueous outflow facility following glaucoma filtering surgery.14,16,18 The potential long-term side effects of hypotony include visual impairment, choroidal detachment, and macular and/or optic disc edema.14,16,19

Several studies have investigated short-term hypotony following AC cannulation or paracentesis without the ability to measure IOP immediately after needle removal.20–24 Lu et al.9 performed AC paracentesis with a 27-G needle as treatment for elevated IOP in angle closure glaucoma patients; they reported IOP measurements obtained with Goldmann Applanation Tonometry (GAT) prior to AC paracentesis, then 15 minutes and 24 hours after AC needle removal. Intraocular pressure was approximately 30 mm Hg lower than baseline measurement at both 15 minutes and 24 hours after AC needle removal, but no hypotony was reported during the study, as the average baseline IOP was 58 mm Hg, and hence post-paracentesis IOP was approximately 28 mm Hg.6 Several other studies report similar results, also showing the safety and efficacy of AC paracentesis for the treatment of elevated IOP associated with acute angle closure.7–9 These results don’t represent normal ocular behavior or hypotony recovery times however, as the study patients had closed iridocorneal angles that limited aqueous outflow facility.

Gerometta et al.25 performed AC cannulation and paracentesis with a 28-G needle and recorded recovery times
following varying aqueous humor volume withdrawals from the AC. Intraocular pressure was measured with a Perkins tonometer in sheep and Tono-Pen in rabbits. In sheep, 60, 120, and 300 μl of aqueous humor was withdrawn in different animals, and they reported recovery times to baseline IOP of 49, 56, and 50 minutes, respectively. Whereas in rabbits, a 13-minute recovery time to baseline IOP levels was reported after 50 and 100 μl of aqueous humor was withdrawn. All these studies were hampered by the lack of a reliable IOP measurement technique at low IOPs, as well as intermittent snapshot IOP measurements that fail to capture the time course of recovery from hypotony. Studies of AC paracentesis to acutely lower IOP in CRAO report visual acuity as the outcome variable, but do not address hypotony or IOP recovery times from hypotony following AC cannulation. Furthermore, there are no studies characterizing the ability of the eye to recover from cannulation-induced hypotony after repeated AC cannulations.

Previous studies have reported intermittent IOP measurements recorded with tonometers that have 1 to 3 mm Hg inherent measurement error, to characterize IOP no sooner than 15 minutes post cannulation. We have collected true magnitudes of hypotony after repeated AC cannulations. Furthermore, there are no studies characterizing the ability of the eye to recover from cannulation-induced hypotony after repeated AC cannulations.

METHODS

Animals

All animals were treated in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research under a protocol approved and monitored by the Institutional Animal Care and Use Committee of the University of Alabama at Birmingham. Seven eyes from four male rhesus macaques, aged 3- to 6-years old, with no ocular abnormalities were used for data collection for this study, which was conducted as standard procedure within a larger National Institutes of Health (NIH)-funded study aimed at determining the contribution of IOP fluctuations to glaucoma onset and progression. All animals were kept on a 6 AM to 6 PM light-dark cycle and fed at approximately 6 AM and 2 PM daily. All animals received water through a continuous feed that was available at all times. Food and water intake was not measured for this study. Ketamine (3 mg/kg) with dexmedetomidine (50 mcg/kg) used as the induction anesthetic for all experiments, followed by isoflurane inhalant anesthesia (1%-3%) for maintenance during AC cannulation. All NHPs were kept warm with a warming blanket and systemically monitored for heart rate, SpO2, end tidal CO2 volume, electrocardiogram (EKG), and temperature with documentation every 15 minutes during all procedures. All eyes were prepped with 5% betadine solution applied to the eye and lids, followed by a double rinse with sterile balanced salt solution prior to AC cannulation. Three drops of 2% proparacaine topical anesthetic were then instilled to minimize discomfort. Polymycin B antibiotic ointment was applied to all eyes following needle removal. There were no signs of either discomfort (observation of NHP behavior) or persistent ocular inflammation (follow-up slit-lamp exam) following bilateral AC cannulation needle removal, even after 10 sessions.

Bilateral IOP Telemetry System

We have developed and validated an implantable telemetry system that wirelessly records 500 measurements of IOP per second for up to 2.5 years. The study protocol mandates a minimum 4-week recovery time following surgery. Using an enhanced version of this system, continuous bilateral IOP, bilateral electro-oculogram (EOG), and aortic blood pressure were recorded both before and after AC cannulation with a 27-G needle at the corneal limbal junction in 10 different sessions, 2 weeks apart in each eye of the NHPs. The IOP transducers were calibrated via AC manometry, and all data corrected for signal drift. Bilateral IOP was recorded continuously while the NHPs were awake and behaving preprocedure and data collection, under general anesthesia during IOP transducer calibration experiments, then continuously after AC needle removal was removed until IOP recovered to baseline levels. The animals were given an anesthesia reversal agent (antisedan) soon after AC needle removal, and hence were not kept anesthetized for the recovery of IOP to 6 and 10 mm Hg. Thus, the impact of anesthesia on IOP recovery was minimized.

Precannulation baseline IOP was calculated by averaging 60 seconds of continuous telemetric IOP data for each eye following anesthetic induction. Postcannulation IOP after removal of needle from the AC was calculated by averaging 30 seconds of continuous telemetric IOP data for each eye. The mean IOP for each time point was used for analysis. The times necessary for IOP to recover from post-cannulation levels to 6 and 10 mm Hg, respectively, were calculated. The data were collected every 2 weeks over a 5-month period for a total of 10 sessions per eye in each NHP.

Statistical Analysis

Separate linear regression models were used to assess the linear correlation between magnitude of hypotony, and in recovery times to 6 and 10 mm Hg over the 10 sessions. Statistically significant differences were not observed, so data from each eye over 10 sessions were averaged and summarized as means and SD. A Wilcoxon signed-rank test was used to determine if the change in IOP from baseline to post-AC cannulation was significantly different from zero to test the hypothesis that significant hypotony is present after AC cannulation needle removal, with statistical significance defined as P less than 0.05.

RESULTS

The Table shows the baseline IOP, IOP immediately following AC needle removal, the magnitude of IOP change from baseline to AC needle removal, and the time for IOP to recover to 6 and

<table>
<thead>
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<th>Parameter</th>
<th>Mean</th>
<th>SD</th>
<th>Minimum</th>
<th>Maximum</th>
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<td>IOP at baseline, mm Hg</td>
<td>11.5</td>
<td>1.5</td>
<td>9.7</td>
<td>15.2</td>
</tr>
<tr>
<td>IOP after AC needle removal, mm Hg</td>
<td>2.3</td>
<td>0.4</td>
<td>1.7</td>
<td>3.5</td>
</tr>
<tr>
<td>Magnitude of IOP change, mm Hg</td>
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<td>1.5</td>
<td>−10.7</td>
<td>−6.7</td>
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<tr>
<td>Time to 6 mm Hg, min</td>
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<td>4</td>
<td>27.2</td>
<td>39.8</td>
</tr>
<tr>
<td>Time to 10 mm Hg, min</td>
<td>63.7</td>
<td>15.2</td>
<td>43.7</td>
<td>92.1</td>
</tr>
</tbody>
</table>

The Table shows the baseline IOP IOP immediately following AC needle removal, the magnitude of IOP change from baseline to AC needle removal, and the time for IOP to recover to 6 and.
10 mm Hg after AC needle removal in seven eyes of four NHPs. The magnitude of IOP change significantly decreased from baseline to immediately following AC needle removal ($P = 0.0156$). Figure 1 shows the mean IOP at baseline and immediately after AC needle removal in each eye. As shown, the IOP at baseline and after AC needle removal was similar for all eyes, ranging from 9.7 to 13.2 mm Hg at baseline, and 1.7 to 3 mm Hg after AC needle removal. There was no significant change in IOP between sessions within eyes ($P = 0.58$). Figure 2 shows the mean recovery time by NHP to 6 and 10 mm Hg. The mean recovery time to 6 mm Hg was 52.4 minutes (SD 4.0), ranging from 27 to 40 minutes. The mean recovery time to 10 mm Hg was 63.7 minutes (SD 15.2), ranging from 44 to 92 minutes.

Over the course of 10 AC cannulations carried out every 2 weeks for 5 months in all eyes, the magnitude of hypotony ($P = 0.58$), the time course of recovery to 6 mm Hg ($P = 0.15$), and the time course of recovery to 10 mm Hg ($P = 0.68$) were not significantly different with each session (Figs. 3A–C). Figure 3A shows the IOP immediately following AC needle removal by eye in each NHP by session.

**DISCUSSION**

We used continuous IOP telemetry to measure both the level of hypotony and the precise time needed for IOP to recover to 6- and 10-mm Hg bilaterally in each NHP after AC cannulation, except for one NHP due to unilateral telemetry. Results show that there is significant hypotony immediately following AC cannulation with a 27-G needle, but IOP recovers to physiologic levels (above 6 mm Hg) within approximately 30 minutes after needle removal.

Previous animal studies did not record postoperative or post-procedure IOP measurements until at least 15 minutes after AC needle removal, which was not quick enough to record the hypotonous period measured in our study. In studies of AC cannulation and paracentesis in sheep and rabbits, IOP recovery was not assessed using accurate continuous measurement; although IOP recovered within an hour.$^{23}$ These recovery times are somewhat similar in duration to our measurements.

Carnahan et al.$^7$ performed AC paracentesis in series in five cases of acute elevation of IOP reporting the safety of repeated AC paracentesis with low risk to the patients. Our study demonstrates that IOP recovers quickly to physiologic levels after AC cannulation. In addition, IOP recovery time following AC cannulation every 2 weeks remained consistent over a 5-month period. Hence, repeated AC cannulations do not result in persistent corneal wound leaks or persistent hypotony over long study periods. These data also inform clinical care, as previous studies showed no hypotony 1 day after procedures using a 27-G needle. Given the similarities between human and
NHP eyes, postoperative hypotony is likely to occur in these patients, but is likely of short duration.

The present study limited by the following considerations. While the telemetric IOP measurements reported herein are accurate to within \( \pm 0.2 \) mm Hg\(^{32}\) and allow for continuous IOP monitoring over long periods, the NHPs were anesthetized during both the AC cannulation procedure and the follow-up period in which we assessed IOP recovery. Anesthesia could affect aqueous production and/or outflow rates, which may affect the rate of ocular volume recovery, and hence the magnitude and duration of hypotony. There are some differences in aqueous humor dynamics in humans and NHPs, so this result may not be directly translatable to clinical applications.\(^{33–36}\) Also, IOP fluctuates significantly with blinks and saccades,\(^{35,37}\) and it is possible that these IOP transients could prolong AC cannulation wound leakage, and therefore prolong the period of hypotony associated with these procedures in awake behaving NHPs. For the purposes of this

**Figure 3.** (A) Intraocular pressure immediately following AC needle removal by eye in each NHP by session. (B) Recovery time to 6 mm Hg after AC needle removal by eye in each NHP by session; (C) recovery time to 10 mm Hg after AC needle removal by eye in each NHP by session. OD, right eye; OS, left eye.
study, the results and conclusions remain valid for NHPs, and may translate to humans.

Anterior chamber cannulation is common in animal studies and human patient procedures, but its lasting effects on IOP are unknown. Results show that the IOP falls to very low levels after AC needle removal; the eye is only hypotonous for approximately 30 minutes after needle removal, and recovers to near baseline levels within approximately 65 minutes. Most importantly, repeated AC cannulations have no effect on either the magnitude of hypotony or the recovery time to either 6 or 10 mm Hg, which is very important for studies involving repeat AC cannulations. This should be taken into account in research or clinical settings when consideration of IOP or its effects is important, but results show that IOP recovers quickly in NHPs even after repeated AC cannulation procedures in the same eye.

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