# The pressure and volume dependence of the rate of wash-out in the bovine eye* 

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## ABSTRACT

The rate of increase of outflow facility (the wash-out rate) was measured in bovine eyes at 6 and 15 mm Hg . The time-rate-of-change of facility was less at $6 \mathrm{~mm} \mathrm{Hg}(0.20$ : $\Delta$ facility/hour) than at $15 \mathrm{~mm} \mathrm{Hg}(0.44)$. However, when the data was analyzed as a function of volume passing through the outflow system, the volume-rate-of-change of facility was the same at $6(0.35: \Delta$ facility $/ \mathrm{ml})$ and $15 \mathrm{~mm} \mathrm{Hg}(0.34)$. This was consistent with the hypothesis of macromolecules "washingout" of the aqueous outflow system, if these macromolecules were saturable in the perfusate.

When aqueous humor outflow facility is determined by perfusion methods in live (1) or enucleated eyes (non-human) (2), a steady increase in measured outflow facility with time is observed. This increase is known as the "wash-out effect". The cause of this flow-induced facility increase has not been established but has been postulated to result from a "washingout" of extracellular material (2) (such as glycosaminoglycans (3) or proteins (4)) from the aqueous outflow pathway. Recent studies, however, have shown minimal evidence of "wash-out" of glycosaminoglycans (5). In this study, we have examined the wash-out effect at two different perfusion pressures (6 and 15 mm Hg ), and have investigated the relationship between the wash-out rate and the volume of perfusate passing through the aqueous outflow pathway. Our results are consistent with the hypothesis of macromolecules "washing-out" of the aqueous outflow pathway.
Enucleated bovine eyes from two-week old calves were obtained from a local abattoir (Arenas \& Sons, Hopkinton, MA) and transported in ice-saline. Perfusions were performed, beginning within eight hours postmortem, using Dulbecco's phosphate buffered saline (Life Science Technologies Inc., Chagrin Falls, OH) with 5.5 mM glucose added. A total of 21 eyes were perfused at either 6 or 15 mm Hg using a computer controlled syringe pump (Model 944, Harvard Apparatus Co., South Natick, MA). The eyes were placed into a gauze-filled
beaker, wetted until the cornea was just covered with saline, and then placed into a constant temperature bath set to a temperature of approximately $33^{\circ} \mathrm{C}$. Anterior chamber deepening was prevented by placing the perfusion needle in the posterior chamber or by placing an iridotomy in the iris.
The experiments were conducted such that the perfusions at 6 mm Hg were typically longer ( $4-8$ hours) than those at 15 mm Hg (3-4 hours) in an attempt to perfuse the same total volume of fluid through the eyes. Typically $2-4 \mathrm{ml}$ of perfusion fluid passed through the outflow pathway of the low pressure eyes while 3-5 ml passed through the high pressure eyes. Pressure and flow were monitored continuously throughout the experiment.
For data analysis, an average facility was determined for sequential time periods that varied between 2 and 5 minutes, depending on the total time of perfusion. The initial data during the equilibration period ( $30-120$ minutes at 6 mm Hg ; $10-45$ minutes at 15 mm Hg ) were rejected. The facility data were then fitted, either as a function of time ( $t$ ) or volume that had passed through the perfusion system (V), as

$$
c(t)=a_{0}+a_{1} t+a_{2} t^{2}
$$

or

$$
c(\mathrm{~V})=b_{0}+b_{1} \mathrm{~V}+b_{2} \mathrm{~V}^{2}
$$

with $a_{1}$ and $b_{1}$ the initial slopes representing the time or volume rates of wash-out, respectively. Correlation coefficients ( r ) were always greater than 0.97 and usually greater than 0.99 .
Data from 3 eyes were rejected as too noisy to get a reliable slope, from one eye because of pump error, and from one eye $(6 \mathrm{~mm} \mathrm{Hg})$ because its initial slopes ( $a_{1}$ and $b_{1}$ ) were 4 standard deviations different from the mean value of that group (in fact, this eye had a wash-out rate that was greater even than any eye in the 15 mm Hg group).

Typical results are shown in Fig. 1 comparing a pair of eyes,

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Figure 1: Outflow facility in a pair of eyes as a function of time or as a function of volume of perfusion fluid passing through the aqueous outflow system.
perfused at 6 and 15 mm Hg respectively, as to their wash-out rates as a function of time and of total volume of perfusate that had passed through the outflow pathway. The combined results for the second set of eyes are shown in Table 1: While the time-rate-of-change of outflow facility was different at the two different perfusion pressures, the volume-rate-ofchange of outflow facility was actually the same.
These experiments were motivated by the observation that there appeared to be little or no "wash-out" at a perfusion pressure of 6 mm Hg . As this is near the physiological pressure drop across the aqueous outflow pathway, we queried whether "wash-out" might represent, at least in part, an artifact of perfusion experiments performed at higher and thus nonphysiological pressure drops across the aqueous outflow system..We noted that eyes perfused at lower pressure have less total perfusate passing through the system, and thus we examined what the relationship was between the increase of outflow facility and the quantity of perfusate that passed through the outflow pathway. We found that the volume-rate-of-change of facility was nearly identical at the two pressures examined even through their time-rate-of-change was markedly different.
These results suggest that the "wash-out effect" may not be due to degenerative changes since these would lead to changes proportional to time, not volume of fluid perfused. This type of experiment might therefore provide important insights into the normal functioning of the aqueous outflow system. Our results are more consistent with the notion that some resistancecausing macromolecule may be "washing-out" from the outflow pathway. If so, the concentration of these resistance-

Table 1: Wash-out rate as a function of time and perfusion volume.

| Perfus. <br> Press. <br> $(\mathrm{mmHg})$ | Initial <br> Facility <br> $(\mu \mathrm{L} / \mathrm{min} / \mathrm{mm}$ <br> $\left(\mathrm{a}_{0}, \mathrm{~b}_{0}\right)$ | $\mathrm{dc} / \mathrm{dt}$ <br> $(\mathrm{c} / \mathrm{hour})$ <br> $\left(\mathrm{a}_{1}\right)$ | $\mathrm{dc} / \mathrm{dv}$ <br> $(\mathrm{c} / \mathrm{ml})$ <br> $\left(\mathrm{b}_{1}\right)$ |
| :--- | :---: | :---: | :---: |
| 6 <br> $(\mathrm{~N}=8)$ | $1.2 \pm 0.08$ | $0.20 \pm 0.046^{*}$ | $0.35 \pm 0.036$ |
| 15 <br> $(\mathrm{~N}=8)$ | $1.3 \pm 0.10$ | $0.44 \pm 0.046^{*}$ | $0.34 \pm 0.033$ |

* Difference statistically significant with $\mathrm{p}<0.005$ (two-sided t test)
causing macromolecules in the perfusate must be fairly constant since the rate of flow appears to determine the rate of "washout". Accordingly, increasing flow would increase the net flux of material, thereby reducing outflow resistance. Regional differences in outflow might be expected to amplify the rate of resistance decrease.
The results further implicate a mechanism that causes the concentration of resistance-causing substances in the perfusate to be independent of volume flow rate. This would occur if that substance were readily saturable in the perfusate (such as a relatively insoluble macromolecule).
These observations suggest that a "washing-out" of macromolecules from the outflow pathway might be involved in regulation of aqueous humor outflow resistance. Why this same phenomenon does not occur in the human eye (7) remains a puzzling difference.


## ACKNOWLEDGEMENTS

This study was supported by NIH grants EY05503 and EY01894 and a grant from National Glaucoma Research, a program of the American Health Assistance Foundation.

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*This work was presented in part at the annual meeting of the Association for Research in Vision and Ophthalmology, 1990, Sarasota, Florida.

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[^0]:    Received on January 11, 1991; accepted on March 18, 1991

