# ATTEMPTS TO PROLONG AND INTENSIFY SPINAL ANESTHESIA BY THE ADDITION OF EPHEDRINE, NEOSYNEPHRIN OR EPINEPHRINE TO A PONTOCAINE-GLUCOSE SOLDTION * 

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Vanious substances have been added to spinal anesthetic agents in an attempt to prolong their action. Starch, gliadin and human plasma have all been used for this purpose. More popular, howeser, are the drugs which raise blood pressure-ephedrine, neosynephrin, methedrine and epinephrine having been studied. It was assumed that these drugs might produce the desired result through local vasoconstriction and prevention of vascular absorption of the anesthetic from the subarachnoid space. It has also been suggested that the pressor drugs themselves may, through a direct action on nerve roots, produce partial degrees of block of conduction in these roots (1,2). Another possibility is that the pressor drugs might exert an antagonistic action against those enzymes which destroy local anesthetic drugs. There is no evidence at the moment for this postulate.

A few studies on animals and many clinical reports have appeared recently, indicating (a) that the duration of anesthesia is prolonged to varying degrees by different pressor drugs; (b) that the effective dose of a spinal anesthetic agent can be reduced by such a combination, and (c) that ephedrine, at least, may be able to produce nerve block in its own right (2-6).

We have attempted to assess the value of such a combination by analyzing statistically any change in the increased duration of sensory and motor block which resulted from the addition of a pressor drug. Ephedrine, neosynephrin and epinephrine have been studied.

## Method

A standard dose of pontocaine hydrochloride was administered to each of 539 patients. Ten milligrams of a 1 per cent solution of the drug, i.e., 1 cc., was used. A standard total volume of 3 cc. was injected in each instance. The remainder of the total volume was made up as noted below.

[^0]As a control group, 105 patients received the above drugs to which was added 1 cc. of spinal flaid withdrawn at the time of the lumbar puncture. Two hundred and eighteen patients received 50 mg . of ephedrine sulfate ( 1 cc. of a 5 per cent solution) added to the pontocaineyrlucose mixture. One hundred and six patients received 1 mg . ( 0.5 ce. of a 0.2 per cent solution) of neosynephrin hydrochloride and 110 were given 0.5 mg . ( 0.5 cc. of a 0.1 per cent solution) of epinephrine. In the latter two groups 0.5 cc . of spinal fluid was added to bring the total volume to 3 cc . (table 1 ).

TABLE 1

| No. of Cason | Pontocaine 1\%. | Gluegee 10\%. | Premar Drus | $\underset{\text { ce. }}{\text { Spinel Fiuid. }}$ | Total Volume. |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 105 | 10 | 1 | None | 1 | 3 |
| 218 | 10 | 1 | Ephedrine ( 50 mg ) | None | 3 |
| 106 | 10 | 1 | Vecsynephrin | 0.5 | 3 |
| 110 | 10 | 1 | Fpinephrine ( 0.5 mg .) | 0.5 | 3 |

Lumbar puncture was performed with a 20 -gauge stileted needle, introduced into the third or fourth lumbar interspace. A pressor drug was usually administered intramuscularly prior to the lumbar puncture. The amount and type of this substance varied throughout the series.

TABLE 2
Number of Cases

|  | Control | Ephedrine | Neomynephrin | Epinephrine |
| :--- | :---: | :---: | :---: | :---: |
| 1. Age, years |  |  |  |  |
| $0-10$ | 3 | 18 | 3 | 5 |
| $20-29$ | 13 | 27 | 12 | 16 |
| $30-39$ | 15 | 44 | 18 | 14 |
| $40-40$ | 29 | 54 | 19 | 23 |
| $50-59$ | 23 | 47 | 26 | 37 |
| $60-69$ | 15 | 19 | 24 | 0 |
| $70-79$ | 5 | 7 | 4 | 5 |
| S0-89 | 2 | 2 | 0 | 1 |
|  |  |  |  |  |
| 2. Sex | 60 | 117 | 48 |  |
| Male | 45 | 101 | 58 | 50 |
| Female |  |  |  |  |
| 3. Region of Operation | 23 | 61 | 24 |  |
| Upper Abdomen | 25 | 45 | 28 | 23 |
| Lower Abdomen | 26 | 54 | 26 | 42 |
| Inguinal |  |  |  | 28 |
| Below Inguinal |  |  |  |  |

After injection of the anesthetic mixture the sensory level to pin prick was recorded at five to fifteen minute intervals during the operation and at thirty minute intervals after the operation until the level had receded to the twelfth thoracic or first lumbar segment. The dermatome classification of Foerster was used.

During the postoperative period patients were urged to move any portion of their lower extremities and the first voluntary movement was recorded as the end point of motor paralysis This might have been motion of a toe, or motion of the whole extremity.

Patients in the series were subjected to a variety of sargical procedures below the diaphragm. They ranged in age from 13 to 88 years; in weight from 97 to 270 pounds, and in height from 60 to 74 inches (table 2).

## Results

A. Duration of Motor Paralysis.-Motor paralysis of the lower extremity was considered as being present from the time of injection of the anesthetic agent until the first voluntary movement of any part of the lower extremity. Table 3 indicates the values found for the control series and for the groups given various vasoconstrictor drugs.

It is apparent that the addition of 50 mg . of ephedrine sulfate to the pontocaine-glucose mixture did not change the duration of motor block as we have defined it except in the perineal and lower extremity group. Neosynephrin and epinephrine in the dosages used prolong motor paralysis in all groups. The increased duration of block following these drugs was statistically significant. There was no essential difference between the two drugs from the standpoint of their ability to increase the time of motor block.
B. Duration of Sensory Anesthesia.-Sensory anesthesia was measured from the time of injection of the anesthetic misture until the sensory level to pin prick had receded to the twelfth thoracic dermatome in all operations except those below the inguinal area; these were measured to the first lumbar. Table 4 lists the duration of anesthesia for the control and vasoconstrictor groups.

Again it is evident that ephedrine did not inerease the duration of sensory anesthesia produced by the dose of pontocaine used in this series. Neosynephrin and epinephrine did increase the length of sensory block significantly according to statistical analysis. The difference in action of the two drugs was not significant.
C. Rate of Decline of the Sensory Level.-In an attempt to analyze the minute to minute effect of the addition of pressor drugs, graphs of those cases in which the sensory level of anesthesia reached or exceeded the fifth thoracic dermatome were constructed. The actual sensory level at fifteen to thirty minute intervals was charted for each patient (fig. 1).

TABLE 3
Duration of Motor Block from Time op Injection
until First Movement of Lower Extremity

| Ilcgion | Control |  |  | Ephodrine |  |  | Nocoynephrin |  |  | Epinephrine |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Duration | Minutem | Caven | Minutes | Cases | $\left\lvert\, \begin{gathered} \text { Per } \\ \text { Cent } \\ \text { Increase } \end{gathered}\right.$ | Minutes | Cases |  | Minutea | Cases |  |
| Upper Abdomen | Average Median Range | $\begin{aligned} & 175 \\ & 170 \\ & 105-270 \end{aligned}$ | 22 | $\left\|\begin{array}{l} 195 \\ 185 \\ 115-330 \end{array}\right\|$ | 53 | 11 | $\left\|\begin{array}{l} 232.5 \\ 230 \\ 150-315 \end{array}\right\|$ | 20 | 32** | $\left\|\begin{array}{l} 259 \\ 255 \\ 195-3-45 \end{array}\right\|$ | 20 | 48*** |
| Lower Aladomen | Average <br> Median <br> Range | $\left\|\begin{array}{l} 195 \\ 192.5 \\ 95-270 \end{array}\right\|$ | 22 | $\left\|\begin{array}{l} 198 \\ 197.5 \\ 130-315 \end{array}\right\|$ | 42 | - | $\left\|\begin{array}{l} 276 \\ 265 \\ 180-410 \end{array}\right\|$ | 20 | 41** | 276 <br> 275 <br> 140-390 | 36 | 41** |
| Inguinal | Average <br> Median <br> Range | $\left\|\begin{array}{c} 180 \\ 175 \\ 60-295 \end{array}\right\|$ | 25 | $\left\|\begin{array}{l} 205 \\ 205 \\ 100-350 \end{array}\right\|$ | 46 | 13 | $\left\|\begin{array}{l} 255 \\ 245 \\ 155-495 \end{array}\right\|$ | 23 | 41** | 254 255 190-340 | 11 | 41** |
| Perincal and Laver Ext. | Average Median Range | $\begin{aligned} & 149 \\ & 150 \\ & 65-260 \end{aligned}$ | 27 | $\left\|\begin{array}{l} 196.5 \\ 190 \\ 75-315 \end{array}\right\|$ | 48 | $31^{* *}$ | $\left.\begin{aligned} & 245 \\ & 250 \\ & 85-400 \end{aligned} \right\rvert\,$ | 22 | 6-** | $\left\|\begin{array}{l} 248 \\ 252.5 \\ 135-350 \end{array}\right\|$ | 24 | 66** |
| Total Average |  | 173 | 96 | 198 | 189 | 14 | 252 | 85 | 45** | 262 | 01 | 51** |

[^1]TABLE 4
Duration of Sensory Block prom Time of Injection until Sensory Level Descended to $\mathrm{D}_{12}$ or $\mathrm{L}_{1}$ *

| Rerion | Control |  |  | Ephedrine |  |  | Neomsaephrin |  |  | Epinephrine |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Duration | Minutes | Cases | Minutes | Cases | $\begin{gathered} \text { Per } \\ \text { Cent } \end{gathered}$ | Minutes | Cases | $\left\|\begin{array}{c} \text { Per } \\ \text { Cent } \\ \text { Incrense } \end{array}\right\|$ | Minutea | Cases | $\underset{\text { Increaso }}{\text { Per }}$ |
| L'pper Abrlonuen | Average Median Range | $\left\|\begin{array}{l} 244 \\ 240 \\ 130-160 \end{array}\right\|$ | 22 | $\begin{aligned} & 238 \\ & 240 \\ & 150-315 \end{aligned}$ | 40 | - | $\left\|\begin{array}{l} 294 \\ 290 \\ 225-340 \end{array}\right\|$ | 18 | 20** | $\left\|\begin{array}{l} 272 \\ 270 \\ 195-330 \end{array}\right\|$ | 19 | 11 |
| Lower Abdomen | Average Median Range | $\left\|\begin{array}{l} 217 \\ 220 \\ 120-300 \end{array}\right\|$ | 22 | $\begin{aligned} & 214.5 \\ & 210 \\ & 135-315 \end{aligned}$ | 32 | - | $\left\|\begin{array}{l} 303 \\ 295 \\ 200-395 \end{array}\right\|$ | 18 | 39** | $\left\|\begin{array}{l} 277 \\ 275 \\ 170-390 \end{array}\right\|$ | 37 | 27* |
| Inguinal | Average Median Range | 236 <br> 230 <br> 120-320 | 21 | $\left\|\begin{array}{l} 223 \\ 217 \\ 100-350 \end{array}\right\|$ | 38 | - | $\left\lvert\, \begin{aligned} & 294 \\ & 285 \\ & 185-540 \end{aligned}\right.$ | 19 | 24** | $\left\|\begin{array}{l} 268 \\ 267 \\ 220-340 \end{array}\right\|$ | 12 | 13* |
| Perineal and Extremity | Average Median Range | $\left\lvert\, \begin{aligned} & 180 \\ & 180 \\ & 65-275 \end{aligned}\right.$ | 25 | $\left\|\begin{array}{l} 202 \\ 195 \\ 105-340 \end{array}\right\|$ | 43 | 12 | $\left\|\begin{array}{l} 259 \\ 260 \\ 165-445 \end{array}\right\|$ | 20 | 43** | $\left\|\begin{array}{l} 265 \\ 280 \\ 150-380 \end{array}\right\|$ | 25 | 47** |

[^2]




Fia. 1. Maintenance and rate of decline of the sensory level of spinal anesthesia as affected by the addition of a pressor drug to the anesthetic agent. - Only those cases in which the sensory level reached or exceeded the fifth thoracie dermatome are included.

Much of the information derived from these charts is similar to that described in section B. Certain additional facts are suggested, however. Patients receiving the ephedrine-pontocaine-glucose mixture responded with a higher level of anesthesia than did patients in the other groups. This was associated with a greater tendency for the ephedrine combination to "crawl" in a cephalad direction. It is uncertain whether these observations indicate a delay in fixation of the anesthetic agent, a greater tendency to rise because of the increased specific gravity of the misture or a wider spread of anesthetic effect owing to a specific action of ephedrine itself.

The data suggest that the anesthesia produced by the addition of epinephrine tended to remain longer at the higher levels before beginning to recede. This interpretation of a more intense action is supported by figures listed in table 5.
D. Behavior during Operation.-The incidence of certain subjective complaints and objective findings is given in table 5.

TABLE 5
Incidence of Reactions during Operation

|  | Control | Ephedrine. <br> per cent | Neosynephrin, <br> per cent | Epinephrine; <br> per cent |
| :--- | :---: | :---: | :---: | :---: |
| Emesis | 6 | 8 | 8 | 10 |
| Nausea and retching | 27 | 32 | 30 | 28 |
| Puin | 17 | 8 | 22 | 7 |
| Traction discomfort | 26 | 16 | 20 | 15 |
| Decrease in systolic blood | 26 | 21 | 43 | 40 |
| pressure of 25 per cent or |  |  |  |  |
| more from control systolic |  |  |  |  |
| level |  |  |  |  |

Nausea, retching and vomiting occur frequently during spinal anesthesia. A variety of causes may be listed for this reaction. One of these is the absorption of the local anesthetic agent into the blood stream with subsequent stimulation of a vomiting center in the central nervous system. If a pressor drug delayed rascular absorption of the anesthetic drug from the subarachnoid space it is possible that the incidence of nausea and vomiting might be reduced. The data in table 5 do not support such a contention.

The data in table 5 are of interest as one attempts to evaluate a possible change in the intensity of anesthesia produced by the addition of a pressor drug. The more complete the block of nerve roots the lower should be the incidence of pain experienced by the patient during operation. These data indicate that although the increased duration of block is about equal with neosynephrin and epinephrine, there is a more "solid" type of anesthesia produced by the addition of the latter drug. Ephedrine likewise appears to increase the intensity of the anesthesia if one can so interpret the figures for pain and traction discom-
fort listed in the table. Confirming the observations of others $(3,4)$ no evidence of any systemic action of the sympathomimetic drags was found. The incidence of hypotension during anesthesia was, as a matter of fact, greater in the groups receiving neosynephrin and epinephrine.
E. Postoperative Complications.-All patients in the series were carefully observed during their stay in the hospital following operation. The incidence of the more common postoperative sequelae is presented in table 6.

TABLE 6
Inctbence of Postoperative Sequelae

|  | Control | Ephedrine. per cent | Neonynephrin, per cens | Epinephrine, per cent |
| :---: | :---: | :---: | :---: | :---: |
| Nausca and emesis | 27 | 21 | 12 | 20 |
| Urinary retention | 15 | 13 | 8 | 11 |
| Atelectasis and bronchopneumonia | 4 | 8 | 9 | 3 |
| Pustural headache | 5 | 6 | 5 | 10 |

From these data there is no evidence that the addition of a pressor drug is followed by an increased incidence of untoward effects. There were no instances of the cauda equina syndrome, adhesive arachnoiditis, transverse myelitis or radiculitis in the series.

## Discussion

It should be emphasized that the pontocaine-glucose misture without a pressor drug produced anesthesia which varied widely in duration from patient to patient. Sensory anesthesia ranged from 65 to 320 minutes, motor block from 65 to 295 minutes. Any one patient might therefore respond to pontocaine alone with a block of five hours or more. For this reason, before valid conclusions can be reached as to the ability of a particular pressor drug to increase the duration of anesthesia one must study large numbers of patients and subject the results to statistical analysis.

Some of the data presented in this paper are at variance with opinions expressed by other authors (4-6). No increased duration of action as the result of the addition of 50 mg . of ephedrine sulfate to a standard pontocaine-glucose combination was demonstrated. This does not deny the possibility that the duration of action of a shorter-acting anesthetic agent such as procaine might be increased by ephedrine. Our results suggest, however, that such an increase would probably be insufficient to prove of much clinical value.

Neosynephrin and epinephrine in the amount used did prolong both sensory and motor block, although not to the extent recorded by other
sulted from different amounts of these pressor drugs was not determined. The possibility of there being some "ideal" concentration is real and this concentration may have been far from that used in this study.

Entirely apart from the problem of prolongation of action is the question of an increased intensity of effect. If it can be shown that sympathomimetic drugs produce this result one can either reduce the dose of the anesthetic agent as suggested by some (4), or, with the usual doses of spinal anesthetic agents, can anticipate less discomfort on the part of the patient during surgical manipulation.

Possibilities of harm from the introduction of a vasoconstrictor drug into the subarachnoid space should be considered. Several theoretical objections can be raised. (a) Vasoconstriction might lead to anoxia of nerve tissue through local reduction of blood supply. Reversal of whatever mechanism is responsible for the production of nerve block might therefore be less likely. To date, there has been no evidence of increased neurologic sequelae following such practices. (b) A patient under spinal anesthesia might show extensive muscular block with respiratory inadequacy. In such instances the shorter the action of the anesthetic agent the more rapidly can normal conditions be restored. (c) Muscular activity is alleged to minimize venous thrombosis. Excessive prolongation of spinal anesthesia might, therefore, be followed by an increased incidence of such vascular disturbances. (d) Use of an increased number of ampules in preparation for a spinal anesthesia increases the chance of contamination. This objection, of course, could be corrected by marketing a single ampule.

Although there are no convincing data to confirm these theoretical objections it seems wise to proceed cautiously until more is learned of the nature of the action of pressor drugs in prolonging or intensifying spinal anesthesia. The gain may not be worth the price.

## Conclusions

1. The efficacy of ephedrine, neosynephrin and epinephrine in prolonging and intensifying spinal anesthesia produced by pontocaine has been studied in a controlled series of 539 patients.
2. With the dosage used and under the conditions of the study, ephedrine proved ineffective except in possibly intensifying the action of pontocaine. Neosynephrin and epinephrine were about equally effective in prolonging both sensory and motor block. Epinephrine appeared to maintain a particular initial sensory level longer than neosynephrin. The prolongation of action associated with the use of pressor drugs was less than reported by other workers.
3. No neurologic sequelae were noted in this series.
4. Possibilities of harm from such a practice were discussed. It was suggested that the ultimate use of pressor drugs in the subarachnoid space may depend upon elucidation of the mechanism of action of these agents.

## REFERENCES

1. Schultz, F. H.: Local Anesthetic Properties of Ephedrine Hydrochloride, Aneathesiology 1: 69-71 (July) 1940.
上. Ruben, J. E.; Knmsler, P. M., and Howell, W. Lu.; Jr.: The Spinal Anesthetic Effects of Ephedrine Sulfate: A Preliminary Report, Science 107: 223-294 (Feb.) 1948.
2. Prickett, M. D.; Gross, E. G., and Cullen, S. C.: Spinal Analgesia with Solutions of Procaine and Epinephrine, Anesthesiology 6: 469-475 (Sept.) 1945.
3. Potter, J. K., and Whitacre, Ir. J.: Pontocaine-Dertrose-Ephedrine for Spinal Anesthesia, Ancsthesiology 7: 490-504 (Sept.) 1946.
4. Romberger, F. T.: Spinal Anestbesia-Practical Facts and Common Fallacies-Clinieal Research on Prolonged Spinal Anesthesin Using Vasoconstrictor Adjunctives, Anesth. \& Analg. 22: 252-263 (Sept-Oct.) 1943.
i. Shane, S. M., and Ruiz, E. T.: Use of Adrenalin to Prolong Spinal Anesthesia, Am. J. Surg. 74: 189-191 (Aug.) 1947.

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[^1]:    ** Highly significant statistically (Fishers' t ).

[^2]:    * Significant statistically.
    ** Highly significant statistically

