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THE CONTRACTILE RESPONSE OF THE SPLEEN OF MINIATURE SWINE TO INTRA-ARTERIAL INFUSION OF EPINEPHRINE

BY

LCDR Thomas L. Wachtel, M.D.
CPT G. R. McCahan, Jr., DVM
Mr. William M. McPherson, B.S.

September 1972

U. S. ARMY AEROMEDICAL RESEARCH LABORATORY
Fort Rucker, Alabama 36360

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U. S. Army Medical Research and Development Command

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The Vivarium of the United States Army Aeromedical Research Laboratory is fully accredited by the American Association for Accreditation of Laboratory Animal Care.

In conducting this research, the investigators adhered to the "Guide for Laboratory Animals Facilities and Care" prepared by the committee on the Guide for Laboratory Animals Facilities and Care, National Academy of Sciences, National Research Council. Humane procedures were utilized throughout, and a graduate veterinarian was in constant attendance to perform all surgical procedures and to ensure that all animals were fully anesthetized and insensitive to pain.
ACKNOWLEDGMENTS

The authors are indebted to Janice Speigner, Diana Patrick, John Barbaccia, Tom Downs, Richard Chapman, David Bellemore, Rodney Polk, Malcolm Kirk, C. D. Williams, Patricia Wagner, and Frederick Nelson for their generous assistance, without which this project could not have been completed.
ABSTRACT

The spleen of miniature swine is a blood storage organ which contracts with intra-arterial injection of epinephrine (and presumably other stressful stimuli) and thus autotransfuses the animal. We recommend the removal of the spleen of miniature swine prior to the use of this animal for any shock studies.

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THE CONTRACTILE RESPONSE OF THE SPLEEN OF MINIATURE SWINE TO INTRA-ARTERIAL INFUSION OF EPINEPHRINE

INTRODUCTION

The spleen is the largest lymphoid organ in the animal body and an important reservoir of blood. In the cat and dog, the spleen is a storage organ which autotransfuses the animal and responds to appropriate stimuli; in other animals the spleen is only a rudimentary structure. Despite extensive observations of the pig's liver and spleen during extracorporeal perfusion studies, no data are available from which we might determine the response of the spleen of swine (specifically, miniature swine) to shock-like states.

We anticipate using the pig in a burn shock animal model. These experiments were undertaken to determine whether the contractile response of the spleen was significant when appropriately stimulated and hence whether splenectomies must be performed prior to use of these animals in porcine burn shock studies.

METHODS AND MATERIALS

Twelve (12) white adult male and female Minipigs* weighing an average of 54.7 kg (40.9 to 61.3 kg) were procured, quarantined, and verified to be healthy and free of internal parasites prior to use in this study. They were handled frequently by the vivarium personnel and investigators so that entrance into the pens for studies could be performed without noticeable excitement.

The animals were fasted overnight, premedicated with atropine (1-2 mg) and Innovar-Vet** (1 cc/20 lb), entubated, and anesthetized with Halothane, USP. Central venous and arterial (aortic) catheters were inserted using the method developed in this laboratory. Stainless-steel limb electrodes for electrocardiography were implanted. The animals were allowed to fully recover from the anesthetic and stabilize.

*Modified Pitman Moore Strain of Miniature Swine, Vita Vet Laboratories, Marion, IN 36952
**McNeil Laboratories, Ft. Washington, PA 19304
Each animal was stimulated with an intra-arterial (aortic) injection of epinephrine (3 cc of 1:10000) while the heart sounds were monitored by auscultation and the rhythm evaluated by ECG. Arterial (aortic) blood samples were taken just prior to the stimulation and four (4) minutes after injection of epinephrine. These samples were submitted to study for packed cell volume (PCV), white blood cell count (WBC), platelet count, and peripheral smear. The PCV was obtained by the microhematocrit method. The WBC and platelet counts were obtained using "B-D brand Unopettes*" for dilution, hemocytometer counting chambers, and binocular light microscopy. Blood smears were made and stained with Wright's stain. The stimulation procedure was repeated again before splenectomy. A mean and standard deviation were derived from these determinations.

The Minipigs were premedicated and anesthetized again as described above. The hair over and adjacent to the operative area was closely clipped with a #40 clipper head, washed with a surgical detergent, and prepared with an iodine solution prior to the splenectomy. For the operation, each pig was restrained on its right side. The operative field was draped with sterile towels and Steri-Drape**. Laparotomy was performed through a left subcostal incision. A Bovie Electro-Surgical Unit*** was used for hemostasis and transecting muscle and fat. The spleen was delivered through the incision. The vascular supply to the spleen and the gross anatomy of the spleen were studied. Epinephrine (0.25 to 0.75 cc of 1:0000) was injected into either the splenic or gastro-splenic artery and the response of the spleen was observed. A clean splenectomy and closure of the wound was completed after the method described by Seamer and Walker.7

When the animal had recovered completely the stimulation procedure described above was performed twice. The catheters were removed. Additional blood was obtained by serial superior vena cava sticks for follow-up platelet counts and peripheral smears. A mean and standard deviation were derived from these determinations.

The spleen was submitted for microscopic evaluation. At the completion of the study all animals underwent a complete necropsy.

RESULTS

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*Becton-Dickinson, Rutherford, NJ 07070
**3M Company, St. Paul, MN 55101
***Model CVS, Liebel-Flarsheim Company, Cincinnati, OH 45215
Intra-arterial injection of epinephrine caused an immediate tachycardia (See Figure 1) and frequently premature ventricular contractions. Clinically, the animals were slightly more active and somewhat more irritable.

FIGURE 1. ECG (Lead I) before and immediately after stimulation with epinephrine.
The results of the blood studies are shown in Figure 2. The hematocrit increased an average of 5.43 ± SD 0.94 (or an increase of 13.2% in PCV) with the intra-arterial injection of epinephrine prior to splenectomy and showed no significant increase (0.31 ± SD 0.15%) in PCV after splenectomy (See Figure 2a). The WBC remained unchanged following the intra-arterial injection of epinephrine prior to splenectomy and increased slightly (1828 WBC per ml or a 15% increase) after splenectomy. Platelets were unchanged after stimulation with intra-arterial epinephrine before splenectomy and in the immediate post splenectomy period. There was a transient rise in platelets on the sixth or seventh postoperative day which generally returned to pre-splenectomy levels by the twelfth postoperative day (See Figure 2c). The peripheral smears showed no increase in target cells following splenectomy. There were occasional eosinopenia and lymphopenia.

The spleen of these miniature swine was an oblong, flat, J-shaped organ that measured 3 ± 1 x 6 ± 2 x 35 ± 5 cm and weighed 357 ± 84 gm with blood and 144 gm ± 33 gm with the blood expelled. The spleen had a dual main arterial blood supply (See Figure 3). The larger lateral splenic artery egressed from the pancreatic tissue and often bifurcated prior to entering the curled lateral aspect of the hilus of the spleen. The smaller medial gastro-splenic artery entered the medial one-third of the hilus of the spleen as a large continuing branch of the left gastroepiploic artery. Generally, there was no anatomical anastomosis between these supplies without the spleen. The venous return from the spleen followed the main arterial supply. Vasa brevia also were present between the lateral hilus of the spleen and the fundus of the stomach.

The dual arterial supply was used for studying the contractile response of the spleen to intra-arterial (splenic or gastro-splenic) injections of epinephrine. Small concentrations of epinephrine gave a positive contractile response within a short time and larger concentrations gave a marked splenic contraction (See Figure 4). Only the corresponding medial or lateral half of the spleen contracted in response to the epinephrine indicating a separate arterial supply.

Microscopic evaluation of the spleen showed normal architecture and structures. At necropsy there were a few adhesions to the operative scar in the animals, but no evidence of accessory spleens or splenosis.

DISCUSSION

The spleen is an important reservoir of blood which may be called
FIGURE 2. (A) Mean packed cell volumes (PCV); (B) White blood cell count (WBC); and (C) Platelet count for twelve (12) miniature swine before and after stimulation with epinephrine pre- and post-splenectomy.
injection at arrow

FIGURE 3. Illustration of the spleen of the miniature swine and its vascular supply. Shaded portion indicates contracted spleen from injection of epinephrine into the gastro-splenic artery.
FIGURE 4. Photographs of the spleen of miniature swine before (A) and after (B) injection of epinephrine into the gastro-splenic artery.
upon when the body has a greater need for oxygen in the tissues. This may occur postfeeding, during exercise, following hemorrhage, in carbon monoxide poisoning, during the administration of certain anesthetics, in emotional states, or when the animal is excited, thereby releasing catecholamines such as epinephrine and norepinephrine. These catecholamines cause an increase in blood pressure and the contraction of the spleen, mobilizing erythrocytes into the circulatory system (i.e., autotransfusion). Under these conditions, as well as when exogenous epinephrine is administered in effective doses, there are increased values for erythrocyte counts, packed cell volumes (PCV), and hemoglobin values which may be as high as 15% to 20% in some animals. In an anesthetized dog the values for PCV may be increased from 40% to 45% after a release or injection of epinephrine. Similar considerations presumably apply to the large domestic species, but they have not been demonstrated specifically. We were able to show a significant increase in PCV by intraarterial injection of epinephrine before splenectomy and no significant change in PCV after splenectomy in response to intra-arterial infusion of epinephrine indicating that the spleen of the miniature swine does contract and autotransfuse the animal.

Pre-experiment handling of the animals and our catheterization technique permitted the studies to be done in unanesthetized, unrestrained, non-medicated, stable basal state animals. This minimized the extraneous factors that might have caused splenic contraction. Using only large vessel blood sampling avoided the controversy of large artery and venous PCV being higher than those taken peripherally.

Epinephrine causes myocardial irritability which produces tachycardia and can cause fatal ventricular fibrillation. We used twice the recommended therapeutic intravenous dose of epinephrine and always observed a tachycardia and occasionally arrhythmias. A four-minute post stimulation with epinephrine time was chosen for sampling of arterial blood because of the experience in dogs and because epinephrine is rapidly metabolized (15 to 30 minutes).

Epinephrine causes prompt contraction of the spleen—an effect that is frequently utilized prior to splenectomy [in experimental animals] by injection of the drug into the splenic artery. We demonstrated marked splenic contraction under direct observation (See Figure 4) as our second method of studying the contractile response of the spleen of miniature swine. The dual splenic arterial supply was a useful anatomical finding which enabled us to show segmental contraction of the spleen (See Figures 3 and 4). This configuration is unlike the vascular supply in humans where there is an anastomosis of the left gastroepiploic to a branch of the bifurcated splenic artery prior to entering the spleen. In repeated
examinations and injections of the two arteries, there always appeared to be a distinct anatomical division in the arterial supply.

Several other methods are available for studying splenic contraction. The F_cells factor (PCV body) is influenced by the ability of the spleen (PCV venous) to contract as shown by the fact that it is stable in cats and dogs after splenectomy. We did not use this method because of the inconsistencies in peripheral blood sampling and the inevitable excitability factor of doing so. Roentgenologic observations of the spleen were conducted by Hausner, et. al. He was able to effectively outline the spleen with small, perforated, disk-shaped pellets of lead which were sewed 2 to 3 cm apart along the periphery of the organ. This requires an additional operative technique and stabilization period as well as some restraining technique during roentgenologic studies. Celiac axis catheterization and direct splenoportography each would have considerable technical disadvantages in addition to restraining the animal during roentgenologic examination.

Splenectomy, without excessive blood loss, has no effect on the hematocrit of normal animals. The slight increase in our post-splenectomy PCV is probably the result of contracting one half of each spleen with epinephrine prior to splenectomy; thus autotransfusing the animal to some extent.

The rise in WBC post-splenectomy has been observed by others and its mechanism is poorly understood. Possible mechanisms of this rise in WBC include loss of an inhibiting humoral factor that allows more WBC's to be produced, or the loss of a mechanical filter which then allows the cells once produced to linger longer in the circulation. Likewise, the reason for the increase in WBC's after epinephrine stimulation post-splenectomy in our studies is not clear.

The platelet count data parallel that reported for the rat and approach the transient thrombocytosis seen in humans. No unusual cells were consistently noted on our peripheral smear although Howell-Jolly bodies, Heinz bodies, siderocytes, and target cells are frequently reported.

Gross and microscopic evaluation of the spleens revealed no unusual findings nor did necropsy show any accessory spleens or splenosis which might have confused the blood studies.

CONCLUSIONS
The spleen of the miniature swine has a dual arterial supply.

The spleen of miniature swine contracts with intra-arterial injection of epinephrine.

The spleen of the miniature swine is a storage organ which autotransfuses the animal in response to appropriate stimuli.

We recommend the removal of the spleen of miniature swine prior to the use of this animal for any shock studies.
REFERENCES


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