Use of a novel polyvinyl alcohol membrane as a pericardial substitute reduces adhesion formation and inflammatory response after cardiac reoperation

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Background: Adhesions may increase the incidence of lethal complications of cardiac reoperations, which account for up to 20% of all open-heart surgeries. Herein, we describe the use of a polyvinyl alcohol membrane (PVAM) as a pericardial alternative and describe its performance during reoperation in a relevant animal model.

Methods: The PVAM samples were reticulated by electron beam radiation and manipulated into a tube shape. After thoracotomy, the pericardium of Wistar rats was opened to expose the heart. Rats were treated by pushing the heart back into the thoracic cavity (Sham group), sprinkling the epicardium with talcum powder (Talc group), encircling the heart with PVAM (PVAM group), or sprinkling the epicardium with talcum powder before placing the PVAM to encircle the heart (PVAM + Talc group). Animals were recovered for 8 weeks and then euthanized. Macroscopic findings (ie, extent and severity of adhesions) were classified according to a 4-grade adhesion scale. The PVAM was tested for direct and indirect cytotoxicity with Vero cells. The water absorption capability and in vivo calcification after 8 weeks of subcutaneous implantation of the membrane were examined. Data were analyzed by analysis of variance and Bonferroni post hoc tests.

Results: The PVAM group had lower adhesion scores than the Talc and Sham groups, as well as reduced epicardium thickness and inflammatory cell results, compared with the Talc and PVAM + Talc groups. The PVAM exhibited no direct or indirect cytotoxicity, good water absorption capability (42.4% ± 0.9%), and negligible calcification after 8 weeks (4.42 × 10⁻³ ± 2.56 × 10⁻³ percentage of the total mass).

Conclusions: The PVAM shows promising properties for its potential use as a novel pericardial substitute.

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Pericardial and mediastinal adhesions increase the mortality and morbidity rates after cardiac reoperations.1,2 Many individuals with congenital heart defects will require multiple repeat sternotomies. Numerous strategies have been described for reducing postoperative adhesions, such as sprinkling solutions into the pericardial sac or using expanded polytetrafluoroethylene (ePTFE) or bioabsorbable gelatin sheets.3-5 However, there is no accepted approach for reducing adhesion formation after cardiac surgery.6 Herein, we describe a novel membrane assembled from 10% polyvinyl alcohol (PVA). We examined the use of this PVA hydrogel membrane (PVAM) as a pericardial substitute in an animal model of adhesion formation, focusing on its cytotoxic effects and inflammatory potential.

METHODS
Assembling the PVAM

The PVAM was made as described previously.7 Briefly, the membrane was assembled from an aqueous solution of 10% PVA (Sigma Aldrich, São Paulo, Brazil), which was kept at room temperature for 7 days until dry. The dried membrane was 2-mm thick, and it was manipulated into a tube-like shape (1.5-cm height and 0.5-cm internal diameter).

The PVAM was subjected to a cross-linking step to stabilize the polymer and prevent its dissolution in contacted fluids. This step has prevented PVAM dissolution for at least 24 weeks after implantation in bone tissue.7 Electron beam irradiation, which can simultaneously accomplish the reticulation and sterilization of the membranes, was performed for the cross-linking step. Before irradiation, the material was acetylated by immersion in a solution of 40% formaldehyde, 50% sulfur acid, and 300 g of anhydrous sodium sulfate at 60°C for 24 hours. The PVAMs were washed to remove any residual chemical and stored in distilled water. The membranes were cross-linked by electron beam irradiation with 25 kGy using a Radiation Dynamics electron beam accelerator (Instituto de Pesquisas Energéticas e Nucleares, São Paulo). After this process, the PVAMs were considered to be sterilized and ready for use.
Ethical approval was obtained from the local institutional review board. All animal handling and experiments were performed in accordance with the standards of the Brazilian Council for Animal Experimentation and the 1996 Guide for the Care and Use of Laboratory Animals, published by the National Institutes of Health. Animals were maintained under specific pathogen-free conditions, a 12-hour/12-hour light/dark cycle, and a room temperature of 21°C.

Four-week-old male Wistar rats (average weight, 300 g) were anesthetized with 2% isoflurane. A left thoracotomy was performed, the pericardium was opened, and the heart was exposed. Animals were randomized into 4 groups (10 animals per group). In the Sham group, the heart was pushed back to the thoracic cavity, and the incision was closed. In the Talc group, talcum powder (10 mg) was sprinkled over the epicardium. In the PV AM group, a PV AM tube was inserted into the thoracic cavity and used to encompass the heart from the surrounding tissue, except for where the major vessels exited/entered the heart. Finally, in the PV AM + Talc group, talcum powder (10 mg) was sprinkled over the epicardium, and then a PV AM tube was inserted into the thoracic cavity and used to encompass the heart completely. No sutures were needed to hold the PV AM tube in place, because of its flexibility and the limited nature of the thoracic space.

All animals were recovered from anesthesia for 8 weeks, during which time they were provided standard chow and water ad libitum. Eight weeks after the surgical protocol, the animals were euthanized with an intravenous dose of pentobarbital, the thoracic cavity was opened, and any adhesions were lysed. Two observers (P.P.M.O. and V.P.B.), blinded to each other, scored the adhesions from 0 to 3, according to an adaptation of the score described by Kajihara and colleagues: 0, absence of adhesions; 1, weak adhesions that are easily lysed; 2, moderate adhesions that are lysed with dissection; and 3, severe adhesions that require sectioning by scissors for removal. In all animals, the PV AM tube was inspected for its correct position involving the heart.

Histologic Assessment
Cardiac tissue was fixed in 10% paraformaldehyde, embedded in paraffin, cut into slices, and stained with hematoxylin and eosin. For the inflammatory cell count and epicardial thickness assessments, a midportion of the left ventricle (ie, at the papillary muscle level) was used in all animals. A blinded observer randomly chose 3 optical fields (40× magnification) to evaluate the inflammatory cell count and the epicardial thickness.

Physical and Biological Characteristics of the PVAMs
Indirect and direct cytotoxicity assays were performed with an African green monkey epithelial kidney cell line (Vero cells; Instituto Adolfo Lutz, São Paulo). To assess the indirect cytotoxicity, PVAMs (0.2 g/mL) in the liquid state without cross-linking were mixed in HAM F-10 media (Sigma Aldrich) containing 10% fetal bovine serum (Nutricell, Campinas, Brazil) at 37°C. Different concentrations of the membrane solution were added to cell media containing 3 × 10⁴ cells/mL. After 24 hours, the cell viability was assessed with the (dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide; thiazolyl blue) colorimetric method. Two controls were used: a positive control, containing 10% phenol in the media for 24 hours; and a negative control, containing only media. The hydration capacity of the membranes was evaluated with dry samples of PVAM immersed in distilled water until a stable membrane weight was reached. Five measurements were made, and the hydration capacity was expressed as a percentage. The in vivo calcification index was determined by implanting a PVAM (1 cm²) subcutaneously (s.c.) on the back of the rats. The in vivo calcification index was determined by implanting a PVAM (1 cm²) subcutaneously (s.c.) on the back of the rats.

Abbreviations and Acronyms
CMC = carboxymethylcellulose
ePTFE = expanded polytetrafluoroethylene
PVAM = polyvinyl alcohol membrane

FIGURE 1. Inflammatory cell count (A) and epicardial thickness (B) in all 4 groups after 8 weeks of observation. Bar height indicates mean, with error bar showing standard error of mean. P values by analysis of variance with post hoc Bonferroni for multiple comparisons. PVAM, Polyvinyl alcohol membrane.
the dorsal segment of a Wistar male rat (n = 10). Eight weeks after implantation, the animals were euthanized, and the membrane calcium content was assessed with x-ray fluorescence analysis. The membranes were dried at room temperature and inserted into an x-ray fluorescence spectrometer (EDX 700; Shimadzu, Columbia, Md). The AXIL software package (IAEA Laboratories, Seibersdorf, Austria) was used to determine the calcium content, which was expressed as a percentage of the total mass of the material analyzed.

**Statistical Analysis**
All values are reported as means with SDs. Categorical variables are expressed as frequencies. An analysis of variance with the post hoc Bonferroni test was used for multiple comparisons. Differences with \( P < .05 \) were considered statistically significant. The GraphPad Prism, version 6, software package was used for all analyses (GraphPad Software, La Jolla, Calif).

**RESULTS**
The average membrane hydration capacity was 42.4% ± 0.89%. The average calcium content of PVAMs after 4 weeks of s.c. implantation was \( 4.42 \times 10^{-3} \pm 2.56 \times 10^{-3} \) percentage of the total mass. The number of inflammatory cells was similar between the Sham and PVAM groups and was generally less than the number observed in the Talc and PVAM + Talc groups (Figure 1, A). The epicardial thickness was similar between the Sham, Talc, and PVAM groups and was thinner than the epicardium in the PVAM + Talc group (\( P < .001 \)) (Figure 1, B). No adhesion was observed in any animal in the PVAM group, which was a significant result compared with the other groups (Figure 2). The PVAMs did not demonstrate direct or indirect cytotoxic effects (Figure 3).

**DISCUSSION**
Adhesions offer a major challenge in cardiac surgery reoperations, especially for planned reoperations, such as in patients with congenital heart defects. Various materials have been developed for diminishing pericardial adhesions, such as resorbable collagen membranes, biodegradable collagen-elastin membranes, propylene oxide–treated bovine pericardium, and bioabsorbable gelatin sheets. Herein, we evaluated a novel PVA membrane as a pericardial substitute. In vitro, the PVAM showed a high hydration capacity and no cytotoxic effect after direct or indirect contact with cultured cells. After 8 weeks of s.c. implantation in rats as a pericardial substitute, the PVAMs showed a low calcification index and a low inflammatory effect. In these rats, few adhesions were observed between the epicardium and the surrounding structures of the thoracic cavity. Although PVA-based materials have been used in other biomedical applications, to the best of our knowledge, this report represents the first time that a PVA membrane has been used as a pericardial substitute.

Our findings are consistent regarding the method used, with adhesion scores and microscopic findings obtained...
by a blinded observer. Previous reports have used similar methods with different membranes or biodegradable materials.\textsuperscript{8,14-16} Although we did not compare the PVAM with any commercially available membrane, we found promising results compared with Bel and colleagues,\textsuperscript{17} who reported similar adhesion scores using a sheep model with an absorbable collagen membrane after 8 weeks of implantation. In the present study, we observed almost no adhesions between the PVAM-protected epicardium and the adjacent thoracic structures after 8 weeks of implantation.

A polyvinyl alcohol with carboxymethylcellulose (PVA/CMC) hydrogel has been used in different tissues.\textsuperscript{18,19} Sliker and colleagues\textsuperscript{19} evaluated use of the PVA/CMC hydrogel as an adhesion barrier in colon anastomoses. They found that the hydrogel did not interfere with healing of the anastomotic site and did not prevent adhesion formation. The neutral effect reported by Sliker and colleagues\textsuperscript{19} regarding adhesion formation might be due to the hydrogel format of the PVA/CMC. In particular, the hydrogel was completely absorbed after 3 to 4 weeks and eliminated via urinary excretion. The hydrogel format

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**FIGURE 3.** Results of direct and indirect cytotoxicity tests after 24 hours of cell growth in the 4 groups. Confluence of cells in the negative control (A) and the positive control (B). Confluence of cells under conditions of direct toxicity (C) and indirect toxicity (D). Bar height indicates mean, with error bars showing standard error of mean. \( P \) values by analysis of variance with post hoc Bonferroni for multiple comparisons.
also meant that the precise localization of the PVA/CMC at
the anastomotic site was not ensured. In the present study,
at 8 weeks after its implantation, the tube-shaped PVAM
was not absorbed and continued to maintain its position
encompassing the heart. The cross-linking process used
during membrane assembly ensured that a stable membrane
could be maintained for at least 24 weeks after bone tissue
implantation.

In another study, Slieker and colleagues evaluated use
of the PVA/CMC hydrogel in a rat model of peritonitis, finding
that the PVA/CMC hydrogel reduced adhesions in
the viscera compared with the control group. This result
may be because of the nature of hydrogel, which might
distribute more homogenously between the peritoneum
and abdominal viscera. In a swine model of chronic
ischemia, Lassaletta and colleagues showed that
alcohol use had beneficial effects on precluding adhesion
formation between the pericardium and epicardium.
They simulated coronary ischemia with an aneroid
constrictor to induce chronic coronary insufficiency.
The animals were fed a high-cholesterol diet supple-
mented with either vodka or red wine. The experimental
group that received vodka exhibited fewer pericardial
adhesions, less intramyocardial fibrosis, and reduced
C-reactive protein levels, indicating a lower inflamma-

tory response.

In our study, the PVAM group showed a low inflamma-
tory cell density compared with the other groups. Inflamma-
tory cells play important roles in the healing process and
may increase the presence of adhesions. We used talcum
powder as an irritant agent for better evaluation of the
PVAM properties. Interestingly, we observed that the
PVAM + Talc group showed a similar adhesion score to
the control group. This indirect observation may indicate
that the PVAM protected the surrounding tissues from the
talcum powder that was sprinkled over the epicardium.
Virtually, the only adhesions that did occur were observed
inside the PVAM tube, which was created to simulate the
pericardial membrane (Figure 2).

Our study had some limitations. The model may not
simulate all postoperative and thoracic reentry scenarios
after cardiopulmonary bypass. Talcum was used as an
irritant factor in the absence of blood in the pericardium.
This situation does not reflect the true scenario after an
open-heart surgery. As a result, the rate of adhesion
formation in this model may have been decreased compared
with the clinical situation. Despite the differences
observed among the groups, the period of 8 weeks is short,
and reoperations within 4 weeks are infrequent. Finally, the
mechanism by which PVAM prevented adhesion formation
is not clear and will require further investigation. Despite
these limitations, it was evident that the PVAM elicited little
inflammatory response and precluded the occurrence of
adhesions.

The PVAM offers several clinical advantages, in terms of
its transparency, malleability, elasticity, and possibility of
being sewn into the mediastinum. Some of the commer-
cially available membranes are bovine or porcine derived,
which may elicit an inflammatory response or adhesion
formation. Others, such as those made of ePTFE, are
not transparent and have limited malleability compared with
the PVAM. Our findings offer the possibility of a new
membrane for use as a pericardial substitute before
planned reoperations, especially in newborns and infants
who are more likely to have several surgical interventions
in their pathway for a definitive surgical treatment. This
new membrane may contribute to reduce the occurrence
of adverse events, especially in patients with planned
reintervention because of the nature of their initial disease.

CONCLUSIONS

The use of PVA membranes reduces the occurrence of
adhesions and promotes fewer epicardial reactions, which
may facilitate later reoperation.

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