Embalmimg Techniques for Long-Term Preservation of Bodies

ABSTRACT:
There are only a few methods of preserving bodies that keep them in excellent condition for decades. These methods were developed mostly to sustain the cult of a political leader. Of the long-term embalmed bodies, only V.I. Lenin, Ho Chi Minh, Mao Zedong, Kim Il-sung and Kim Jong-il, the body of Professor Pyrogov, Rosalia Lombardo, and the corpse of a man in the Omsk Anatomical Museum, remain on display. The other bodies were subsequently buried in a tomb. Mention may also be made of the embalmed body of Philippine dictator Ferdinand Marcos, exhibited 1989-2012, which was embalmed by Philippine embalmer Frank Malabed. The body was original with a wax mask on his face. The embalming of the Vatican popes is carried out by the funeral home of Signoracci. Embalming solutions contain formaldehyde, ethanol, glycerine, sodium acetate, potassium acetate and thymol, or use paraffin impregnation. In most cases, it is also necessary to maintain standard environmental conditions such as humidity and air temperature. This paper provides an overview of the major and most successful embalming techniques, with a focus on the Soviet-Russian method and its development.

KEY WORDS: embalming; Lenin; Lombardo; paraffinization; Perón; Pyrogov

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Introduction

The individual components of solutions for long-term embalming each fulfill a certain function. Formaldehyde consolidates proteins and adipose tissue, and inhibits most enzymes and microorganisms, while reducing hemoglobin to methemoglobin. Ethanol stabilizes tissue and converts methemoglobin to pigment, removes epidermal lipids, and increases skin permeability. Glycerine and potassium acetate preserve the pigments and enhance the effects of ethanol. Sodium acetate is used as a pH regulator and preservative. In addition, glycerine maintains tissue elasticity, facilitates distribution of the embalming agent, and draws moisture from the surrounding air to the body. During long-term embalming, i.e. long-term stabilization, water is expelled from the tissues, being gradually replaced with the embalming solution, which results in a hydrothermal equilibrium between the body and the surrounding atmosphere. In most cases, it is also necessary to maintain specific humidity and temperature (Lopukhin, 1997; Frišhons and Vacín, 2014a). Carrying out long-term embalming and research into tissue changes (Fig. 1, Fig. 2) is a very complex process, and for the purposes of this article, the issue has been considerably simplified. With the use of industrially produced solutions, e.g. by Dodge Chemicals (Slocum, 1947), there is no evidence of long-term monitoring of the body for more than a year after the embalming (Labush, 2014).
Materials and Methods

1. The method of D.I. Vyvodtsev was applied in 1881 during the initial embalming of Prof. N.I. Pyrogov's body, using a solution of 5 g thymol, 45 ml ethanol, 2160 ml glycerine, and 1080 ml distilled water (Malyshhev, 1955). Only a few incisions were made; the brain and heart were not removed. The first inspection of the condition of the body by a commission of experts occurred in 1927. In 1940, mold colonies were discovered after opening the coffin. In 1945, the procedure of stabilizing the body, performed by the staff of the Lenin Mausoleum, took 115 days in total. Major preservation work was carried out in 1956 and 1973 by Sinelkov and his team, while in 1979 and 1988 the body was taken to Moscow for additional embalming. Since then, re-embalming has been carried out in 1994, 2000, 2005 and 2011 in a bathtub containing a solution of glycerine, potassium acetate,
and distilled water, with the addition of thymol. This was done in collaboration with the experts from the Moscow Center for Scientific Research and Teaching Methods in Biochemical Technologies, a part of the All-Russian Scientific Research Institute of Medicinal and Aromatic Plants (VILAR). The body was placed in clothing and overalls (Matvejchuk, 2013). Since 2017, the re-embalming has been carried out by a group of Ukrainian scientists, with the participation of Prof. Oleh Melnyk, Prof. Yuri Guminiskii, and their colleagues. The work also included a CT examination, tissue sampling for microscopic scrutiny (Fig. 3, Fig. 4), and other procedures. Meanwhile, the Ukrainian team are developing their own methodology for the regular re-embalming of the body (Fig. 5, Fig. 6) (Hunko, 2019).

2. The method of A.V. Romodanovskii (1904–1969) was used for the embalming of a male body, which took place in the years 1933–1935, at the Institute for Anatomy in Omsk. In addition to freezing, a 10% formalin solution was repeatedly applied through the femoral artery. This process was repeated several times. The body is still stored at room temperature in the Anatomical Museum in Omsk, without periodic treatment. The natural color of the skin and the volume of the tissues are very well preserved (Kuznetsov 1999).

3. The method of Alfredo Salafia (1869–1933), Professor and Sicilian embalmer, who developed his method for the embalming of Rosalia Lombardo in 1920. He applied a solution, which he called the “Salafia perfect fluid”, through the femoral artery: one part glycerine, one part formalin saturated with zinc sulfate and chloride, and one part ethanol saturated with salicylic acid. The body is stored in the Palermo catacombs at a temperature of 25 °C and a relative humidity of 80% (Piombino-Mascali 2006; Piombino-Mascali 2009; Wiesner 2013).

4. The method of Soviet-Russian long-term embalming. Embalming and continuous care for the bodies of V.I. Lenin and other political leaders such as G.M. Dimitrov (1949) (Fig. 7), Kh. Cholbalsan (1952), J.V. Stalin (1953), K. Gottwald (1953) (Fig. 8, Fig. 9, Fig. 10, Fig. 11), Ho Chi Minh (1969), A. Neto (1979), L.F. Burnham (1985), were performed between 1924 and 1991 by experts from the V.I. Lenin Mausoleum Laboratory. Since 1992, the care for Lenin’s body, and the preservation work on the bodies of the North Korean leaders Kim Il-sung (embalmed 1994), Kim Jong-il (embalmed 2012), has been performed by the VILAR. The history of the modern Russian long-term embalming technique dates back to N.F. Melnikov-Razvedenkov (Melnikow-Raswedenkow, 1898). His method was significantly improved by the anatomist V.P. Vorobyov (1876–1937), biochemists B.I. Zbarskii (1885–1954) and S.R. Mardashev (1906–1974), Y.I. Denisov-Nikolskii (1932–2018), V.A. Bykov (1938-) and others. The method consists of the embalming of the body, periodic control and maintenance, and the establishment of supporting technical facilities. Components of the embalming fluid are formaldehyde, ethanol, glycerine, sodium acetate, potassium acetate, and thymol. Mardashev’s experiments on skin samples using the same embalming solution as applied to Lenin’s body determined the evaporation speed of the embalming agent relative to the weight of the sample, thus identifying the optimal ratio of chemicals at a specific temperature and humidity (Lopukhin, 1997).

At the beginning of this method was the embalming of V.I. Lenin, initially performed by Vorobyov, Abrikosov.
and others. A 1%, and then a 2%, formaldehyde solution was first applied to the face, hands, and the front of the body, covered with cotton swabs, as well as injected into certain areas. The body cavities were washed with acetic acid. Subsequently, a 3% formaldehyde solution was applied to the hands and head, and injected into the skull cavity, which had been trepanated. For the whitening of facial and hand tissues, hydrogen peroxide
and ammonia were used after intradermal and muscular application of acetic acid. The whole body was then immersed in a 3% formaldehyde solution for several days. Incisions were made in the abdominal wall, shoulders, forearms, thighs, calves and along the major muscles of the back and in the gluteal region, the skin of the palm, and the lower surfaces of the fingers on both hands. After a week, the body was immersed in a 20% ethanol solution, with the head and hands placed separately in 30 to 35% ethanol, for 6 days. After some time, 20% glycerine was added to the bathtub with 30% ethanol. Following this 14-day ethanol-glycerine bath, the body was immersed in an aqueous solution of glycerine, in May 1924. Eye prostheses were placed in the eye sockets and the eyes were then closed with 2–3 sutures on the edges of the eyelids. The head and hands were regularly soaked in a 1% formaldehyde solution. By the end of June 1924, the body was immersed in 240 l of glycerin, 110 kg of potassium acetate and 150 liters of water, with the addition of a 1–2% solution of quinine and hydrochloric acid. The body was wrapped in rubber elastic bandages, gloves were put on the hands, the body was dressed and placed in a sarcophagus. A petri dish containing an aqueous solution of thymol was placed under the pedestal (Lauer 1924; Melnikov-Razvedenkov 1930; Lopukhin 1997; Zbarskii 1998).

This experimental method has been refined over the decades by further experiments, and the growing experience of the scientists. The resulting methodology is now patented. First, an autopsy is performed to remove the visceral organs while preserving the main blood vessels. This is followed by perfusion of the vascular bed through the arteries with a solution of up to 20% formalin (according to tissue condition, and post mortem interval), with the possible addition of 5% ethanol, 10% glycerin, and 15% potassium acetate. The solution may also be injected into the subcutaneous tissue and muscles. Subsequently, the body is immersed in a bath of the solution for 1 to 2 months. The ratio of body volume to the fixative solution in a triplex silicate glass vessel is 1:10. In the first phase of stabilization of the body by impregnation, a solution is used with a volume of components as follows: glycerin 3–12%, potassium acetate 3–15%, sodium acetate 5%, and up to 100% with distilled water, for 45 days. The wide range of some of the components is because it is an experimental method, and the volume of substances is determined many factors, such as skin pigment, the rate of tissue impregnation, and other variables, known only to the Russian scientists who have been dealing with the issue for decades.

In the second phase, the impregnation lasts 45 days, in a solution consisting of glycerin 12–20%, potassium acetate 15–25%, sodium acetate 5%, up to 100% with distilled water. The third phase lasts 60–75 days, in a solution of glycerin 20–50%, potassium acetate 25–40%, sodium acetate 5%, excipient thymol 0.02% and up to 100% with distilled water. The pH of the solutions must be 8.2–8.4. The ratio of ingredients was determined experimentally, depending on the condition of the body before embalming, skin color, hair color, skin pigment, and subcutaneous fat. Stability of proteolytic enzymes was 64% (Bykov 2002). The procedures are performed by a group of 3 to 5 experts.

The body is stored in a sarcophagus in disposable overalls (jacket, collar, pants, shoe covers, sleeves, drawstrings and drainage valve) made of non-woven hydrophobic material. Sealing tapes at the cuffs and neck are sealed with a 15–30% solution of animal glue gelatin, technical casein, bone glue, and skin glue (Bykov 2013). The overalls are filled with about 10 L of the embalming solution. Clothes are placed on the overalls (Fig. 12). The exact position of the body in the sarcophagus is set by the fixed coordinates (Fig. 13), which ensures perfect adjustment of the lighting system.

![Figure 12. The placement of the body on the lift pedestal in the sarcophagus. Graphics by Kateřina Vacínová.](image-url)
In addition to the technical background for body maintenance, the method also includes daily monitoring of the condition of the body, and possible correction of the humidity of the visible parts of the body, as well as the temperature and humidity of the surrounding atmosphere. Periodic re-embalming after 18 months entails 4 weeks of impregnation in a tank with 200 L of a solution made up according to the condition of the body (Fig. 14). For instance, the body of Georgi Dimitrov was immersed in glycerin 62 liters (31% by weight), potassium acetate 60 kg (30%), and distilled water 78 liters (39%), at a temperature of 14 to 15 °C (CDA, 1957) (Fig. 15). In case of Klement Gottwald, it was a solution of 31% glycerin, 31% potassium acetate and 38% distilled water (National Archive Prague, 1955). The body was weighed before and after impregnation, with an average weight gain of 2 kg. Bonding and sealing of skin incisions was made by rubbing in 15% gelatin dissolved in 10% chloralhydrate. These spots were then covered with silk and another layer of gelatin, and fixed with cotton soaked in 10% formalin. Any soft tissue relief adjustments were made using subcutaneous injections of a liquid mixture of paraffin with white vaseline (8 parts to 2 parts), with wax and gelatin added, at a temperature of 44 °C. The respective spots were then cooled and shaped by hand (National Archive Prague, 1956). Another mixture for correcting the volume of wrinkles and folds is glycerin and potassium acetate 20-40%, in a ratio of 1: 1, thymol 0.02%, hydroxyapatite or bone mineral with a size of 50µ 40-70%, and distilled water up to 100% (Matvejchuk 2019), or Celladamm (a type of dammar resin) 16-20%, white beeswax 55-70%, paraffin...
6-19%, and petroleum jelly 6-12% (Avramov, 2013; Abramov, 2016). Embalming fluid samples were taken for spectrometric examination, and histological specimens were extracted, e.g., from the thighs, for a complex examination of the body’s overall condition. Monitoring and evaluation of the shape and volume of the soft tissues of the head and hands (Fig. 16) were performed at an interval of 4 to 5 years, using reliefometry (a system of relief mapping). As for an assessment of skin color, photoelectric colorimetry was used (Figs. 17, & 18) (Vasilevskii, 1988; Ryabtsev, 1997; Vasilevskii, 2004; Litvinov, 2015; Litvinov, 2016; Matvejchuk 2016). Subsequently, the new results were compared with previous measurements. Moreover, the brightness of the illuminated parts of the body in the sarcophagus, i.e., the head and hands, was measured from the right, left, and front, using a photometer.

Air conditioning system and the technical background of a mausoleum

According to the Soviet-Russian method, the technical facilities of a mausoleum and the disinfection of the laboratory premises are additional preconditions for the successful long-term preservation of embalmed bodies. The equipment of the individual mausolea is similar. Over the decades, improvements have been made to the technology of monitoring and control of the air conditioning system. The facilities consist of a mourning hall with the sarcophagus, a laboratory for observation and body care, and some adjacent spaces. Furthermore, there is a control room and air conditioning machine room. Below, we provide a simplified description of the principles of the air conditioning system in the mausoleum of Klement Gottwald in Prague (early 1960s) and its technical background (Fig. 19). The air conditioning system in the mourning hall was designed to maintain a temperature of 16 °C ± 1 °C and relative humidity of 70% ± 5% in the presence of 80 visitors. The outdoor intake air was filtered and led through the supply channel to the mixing chamber, where it was mixed with the air already discharged from the mourning hall. Through a box dust filter, the air flowed into a pre-heater and a washing device, where it was cooled and saturated to 100% humidity. The air was then led to a fan which drove it into individual distribution pipes with
heaters, which took the air to the required temperature and humidity. The warmed and humidified air entered the mourning hall through the front and side vents. The upper part of the sarcophagus was in a constant stream of air. The air was then sucked out of the mourning hall by a fan separate from the system, and drawn into the mixing chamber, where it was treated again.

Automatic control of temperature and relative humidity was based on a sensor, controlled by a thermostat set to the desired temperature. The automatic temperature and relative humidity control for ventilation and air conditioning in the mourning hall consisted of four parts:

- Thermostatic control of electric outdoor air pre-heater,
- Dew point regulation, signaling, and hall temperature regulation with regulation of the minimum amount of outdoor air.

The sarcophagus in the mourning hall was hermetically sealed. The internal temperature of the sarcophagus was maintained at 16° C ± 1°C under the lights, which was controlled by a remote device with an accuracy of ± 0.1°C. The body was illuminated by a system of 12-volt, 35-watt light tubes with apertures and color filters (Fig. 20). The need for thermal isolation necessitated the placement of dethermal glass under the bulbs, which prevented overheating but did not change the color of the emitted light.

The air conditioning system for the laboratory maintained a temperature of 16° C ± 0.75°C with a relative humidity of 75–85% ± 2.5%. The system contained the same elements, and worked on the same principle, as the system for the mourning hall, except that the air conditioning chambers with pre-heaters were made of sheet metal, and placed upright. Another difference, intended to reduce the occurrence of dust particles and microorganisms, was the inclusion of cotton filters installed on the supply pipe, which cleaned the air before it entered the laboratory. A disinfectant solution was added to the water in the washing device. The walls, ceiling, and floor of the laboratory were lined with washable and disinfectable material. Ethanol (70%) was
used as disinfectant. All equipment had to be made of materials that resisted corrosion and minimized the nutrient surface for microorganisms. A special hydraulic lift with a platform, of a maximum height of 4 meters and a load capacity of 1000 kg, was installed in the laboratory, below the mausoleum. A metal pedestal was mounted on the platform, and a glass casket containing the embalmed body was placed on the pedestal (Fig. 21). An electrically driven, hermetically sealable board was installed in the ceiling. It formed the bottom of the sarcophagus and was closed only when the platform was lowered into the laboratory for a treatment of the body (Fig. 22).

The laboratory’s air conditioning system operated in two modes. In the first, standard mode, the main and backup laboratories were air conditioned in parallel. Ventilation vents and doors in the hermetically sealable wall were opened. Thus, the same temperature and humidity was maintained in both laboratories. The second mode was used for maintenance and additional embalming of the body. The orifices and doors in the airtight wall would be sealed, and the reserve laboratory would be cooled by an evaporator. Only the required temperature would be maintained in the laboratory, while humidity was not controlled. Rooms adjacent to the laboratory were a cloakroom, a doctors’ office, an autoclave room, a lab technicians’ room, and a hallway. Air conditioning of all auxiliary spaces, i.e. the ventilation of the physicians’ office and room for lab technicians, the storeroom, the autoclave room, the ventilation engine rooms, the controller’s room, workshops and toilets, was provided by a separate unit. Air was drawn into the ventilation system of the auxiliary spaces from a common duct, it was then passed through an oil filter, heated, and piped through vents into the individual rooms. An automatically controlled, dry air cooler was installed in the duct for the doctors’ office, lab technicians’ room, storeroom and autoclave room. The air supply to the individual rooms could be regulated by dampers (Frišhons and Vacín 2014b).

5. The paraffinization embalming method was developed by Frédéricq, Hoschetter and Schmeidel, and improved by Professor Pedro Ara Sarría (1891–1973) (Ara, 1934b) who performed the embalming of Eva Perón’s body from July 26, 1952 until November 24, 1955. Initial conservation was carried out with a solution of 10% formaldehyle, 96% ethanol, and 1:1000 solution of water and thymol, injected through the carotid artery, as well as into the visceral cavities and subcutaneous tissue. Subsequently, the body was impregnated with a solution of ethanol, glycerin, potassium acetate, nitrate, and thymol in a 150 L tank. The same solution was used for moistening the face and hands with cotton swabs. The fingers were left wrapped in bandages impregnated with trichlorethylene. Subsequently, the face and hands were treated with cotton swabs soaked in an aqueous solution of 3–5% hydrogen hydroxide. This was followed by a gradual dehydration with 50%, 70%, then 96% ethanol, replacing the tissue water with ethanol. It was later replaced again with a paraffinic organic solvent, turpentine oil, and gasoline, binding the ethanol until the preparation became translucent, which took several
months. Excess paraffin was then removed from the body with the use of filter paper at 56°C. Finally, the body was impregnated in a paraffin tank immersed in a water bath, as well as by vascular infusion. First, paraffin with a low melting point of 36 to 48°C was used, then a higher temperature of 56°C was set using a thermoregulator. This procedure made the solvent evaporate by bubble formation, while the paraffin subsequently hardened in air at room temperature. The body was stored in a sarcophagus with a temperature of up to 25°C. It was protected against mechanical, chemical, and thermal damage, and monitored (Ara 1934a; Ara 1996). Two incisions made to facilitate arterial injection, as well as an incision on the head, were sealed with the impregnating agent. After the repatriation of the body from Spain back to Argentina, it was restored from November 22 to 29, 1974, by Professor Domingo Isaac Tellechea (1935- ) (Tellechea, 1975). The paint covering the neck and shoulders was removed with a solution of petroleum ether. The right shoulder and breasts were restored. A strong insecticidal mixture was applied to a deep incision in the neck, followed by covering the open defects of the neck with hard wax. Synthetic wax was used to replace the small missing part of the left ear. Next, a very fine transparent wax with red cadmium and black pigmentation was applied to adjust the skin color on the head, hands, feet, and the rest of the body. The fingers were repositioned. Lime encrustations and fragments of oxidized iron were removed from the back. At the level of the knee, a wide crack with a large amount of adipose tissue was found. The hair was cleaned and treated. The feet were treated against mold and moisture with an insecticide. The fingernails were treated with a polyester solution in order to conceal the yellowish skin, and to clean and stain the nails. A cast of the missing part of a finger from the right hand was made and put in place. Eyelashes made of natural hair were applied with a needle. Deep defects of the soft tissues of the left anterior shin and right ankle were fixed. An X-ray examination of the body was performed, showing dehydration of the visceral organs. Furthermore, dactyloscopic examination, and a histological examination of the tissue from the right ear was performed (Tellechea 1975). Subsequently, the body was placed in a coffin and interred in a tomb.

Discussion

Preserved embalmed bodies prove that long-term embalming of a human body is possible under certain conditions. Researchers face problems, such as changes in skin color, brought about by changes in melanin due to UV radiation or formaldehyde, which reduces hemoglobin in tissues, and other factors. Changes in tissue volume, drying out, hydrolysis and oxidation of fats, decalcification of bones, or the possible presence of microorganisms in the form of molds or fungi are major problems to deal with. In the initial stages, preservation with formaldehyde, bleaching of brown skin spots with hydrogen peroxide, or treatment of parchment-like spots with water, dilute acetic acid, and hydrogen peroxide is necessary (Lopukhin, 1997). Changes in tissue volume, e.g. swelling, are treated with concentrated ethanol. Constant monitoring and automatic regulation of temperature and humidity in the sarcophagus is necessary (Kozeltsev and Romakov, 2000). Further, non-contact methods for the control of skin condition and color, as well as relief and volume of soft tissues, or the application of advanced imaging methods for monitoring internal structures, have been developed and improved. Last but not least, the continuous analysis of the embalming solution from the overalls, and the collection of tissue specimen to assess the state of cytological and histological structure are of paramount importance. For example, in a specific physiochemical analysis of the skin and epidermis, diffusion parameters of tissue impregnation were found. The histological structure of the skin had been preserved with a number of structural changes in the stratum corneum, cell layers, and dermis, such as disorders of karyolysis, decreased volume of nuclei, decreased DNA content, histones, or RNA concentration. The extent of the changes is determined by the duration of the embalming and the properties of the skin. Substances released during UV absorption, or morphological damage to cells or threshold levels of structural integration of the cell organelle system were also detected (Tomashевич, 1997). A major disadvantage of the Soviet-Russian embalming method is the enormous financial cost of the establishment and operation of the technical facilities.

Postscript

In the 1990s, the staff of the Lenin Mausoleum laboratory studied the effects of the embalming process on the state of epidermal nucleoproteins, using
histochemical detection of nucleic acids and proteins in embalmed materials. They also explored the UV absorbers passing into the fluid during the embalming of skin and muscle tissues, using methods of mathematical physics in order to solve problems with the diffusion of preservative components, or using mathematical models of the process of preservative component diffusion in various tissues, and many other topics (Tomashevich and Nikitina, 1989). No other institution in the world has achieved such knowledge in the study of thanatochemical and microscopic changes in long-term embalmed tissues. Currently, the same team are working on the development of new methods for a non-destructive quantitative assessment of the main components in aqueous solutions containing potassium acetate and glycerin, with the use of Quantitative 1H-NMR spectroscopy (Proton Nuclear Magnetic Resonance) (Abramov 2018; Agrafenin, 2018) or thermogravimetry (Polakov, 2019). While there are a number of research perspectives concerning the long-term preservation of embalmed tissues, it may be assumed that no other laboratory will address these issues at such a level of excellence in the foreseeable future.

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