Severe complication due to inappropriate use of polymethylmethacrylate: a case report and current status in Brazil

Complicação grave do uso irregular do PMMA: relato de caso e a situação brasileira atual

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Introduction: Use of permanent fillers can lead to significant complications. In Brazil, polymethylmethacrylate (PMMA) is a product approved by the Agência Nacional de Vigilância Sanitária (ANVISA), but its use exceeds its indications, leading to serious complications. Recommendations for restricted use have been in place for more than a decade, but cases with serious consequences due to inappropriate use are still seen.

Objective: To report a serious complication due to inappropriate use of PMMA and discuss the current status of PMMA use in Brazil based on recommendations of medical societies and regulatory agencies.

Methods: This report describes a case of extensive necrosis of the gluteal region after injection of PMMA by a non-qualified practitioner; the report also reviews the literature on the current status of PMMA use in Brazil.

Discussion: Despite the efforts of medical societies, acute and chronic complications are still reported in the Brazilian literature. In 2016, more than 17,000 PMMA-related complications were reported; nevertheless, reliable epidemiological data remain unavailable because the number of treatments, the quality of the product, and the training of practitioners remain unregulated.

Conclusion: A significant number of repair procedures are performed in Brazil to correct complications resulting from the use of PMMA. The severity of the reported case highlights the need to combat bad practice by untrained professionals, as well as the need for greater control of PMMA marketing by regulatory agencies.

Keywords: Polymethylmethacrylate; Skin fillers; Brazil; National Health Surveillance Agency; Health advice; Reconstructive surgical procedures/adverse effects.

Abstract

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The most commonly used permanent filler is polymethylmethacrylate (PMMA). PMMA is available as polymeric microspheres ranging in size from 30 to 103 μm. These spheres are suspended in a vehicle of bovine collagen, carboxymethylcellulose, or sodium hyaluronate, which are resorbed after a few days². In 2006, the U.S. Food and Drug Administration approved ArteFill® for use by physicians alone, and limited its application to peri-oral volume augmentation, e.g., nasolabial filling, excluding that of the lips⁴. In Brazil, the National Health Surveillance Agency (Agência Nacional de Vigilância Sanitária, ANVISA) recommends use by trained medical professionals, and does not contraindicate use for body filling or provide guidelines for aesthetic use⁵.

As PMMA is inexpensive and easily obtained, numerous cases of complications have occurred, arising from use by untrained professionals in substandard aesthetic centers. This case report describes complications due to use of PMMA filler in a
clandestine clinic in the city of São Paulo, and discusses the current status of PMMA use in Brazil.

**CASE REPORT**

A 21-year-old female presented with a history of injection of 900 ml of PMMA in the buttocks 12 days prior. The procedure was performed in a beauty salon by a nonmedical professional. She had pain and ulcerated wounds with purulent secretions at the injection site (Figure 1), and had already received ciprofloxacin and clindamycin for 7 days without improvement, as well as day hospital treatment with ceftriaxone and prednisone.

Serial debridement was performed (Figure 1), and continuous negative pressure therapy was applied at 125 mmHg, with sequential exchanges every 3 to 5 days. Suppuration and necrosis of the dermis (Figure 3) and subcutaneous tissue were observed, with nodular formations containing pus and exogenous material in addition to signs of bilateral gluteus maximus fasciitis. Fragments of soft tissue were sent for culture, but no microorganism was identified.

On admission, the patient was afebrile and hemodynamically stable. Laboratory tests showed leukocytosis and elevated C-reactive protein, and computed tomography showed densification and thickening of the skin and subcutaneous tissue of the gluteal regions, in addition to multiple nodular formations consistent with exogenous material/granulomas (Figure 2), but no evidence of collections.

She initially received empirical broad-spectrum antibiotic therapy, but the wound deteriorated.

Figure 1. Surgical wound appearance after serial debridement. a: left lateral, b: posterior, c: right lateral; I: at 14 days; II: after first debridement; III: after second debridement, circumferential increase in surgical wound hyperemia and coalescence of wounds in the sacral region; IV: after third debridement, circumferential increase in surgical wound hyperemia; V: after fourth debridement, appearance of necrosis in the sacral region.

Figure 2. 3D tomographic reconstruction on admission. Blue represents infiltration of subcutaneous tissue in the bilateral gluteal regions at PMMA application sites. A: anterior view; B: posterior view. In retrospective analysis, image of infiltrated area coincided with that of debridement area.

Figure 3. Skin necrosis in a surgical specimen. A: purplish spots on the surface; B: epidermis folded back to show dermis and subcutaneous tissue with extensive necrosis.
Due to the severity of the inflammatory/infectious process, she remained in the intensive care unit for 14 days, and developed acute renal failure. She showed progressive improvement after the 3rd debridement.

Given the size of the wound after sequential debridement and the extensive loss of nutrients, hemoglobin, and microelements, bilateral homologous skin grafting was performed over the gluteus maximus (Figure 4), with addition of negative pressure therapy (Figure 5).

After 3 weeks of treatment with biological dressings (homologous grafts) and improvement of nutritional parameters, autologous skin grafting was performed (Figure 6) with 1:1.5 mesh. The graft showed good integration (Figure 7) without functional deficits, and the patient was discharged after 68 days of hospitalization.

**PMMA in Brazil**

The use of PMMA as a filler has been approved under federal law since 2004 for treatment of HIV-associated lipodystrophy. However, indiscriminate use for aesthetic purposes without scientific evidence merited a public alert by the Federal Medical Council (Conselho Federal de Medicina, CFM) in 2006, as non-qualified practitioners promoted a technique known as “bioplasty”.

In 2007, ANVISA prohibited the preparation of PMMA products by compounding pharmacies in order to regulate quality and purity.

In 2008, complications related to PMMA use in a series of 32 cases led to classification into 5 types: necrosis, granuloma, chronic inflammatory reaction, lip complications, and infection. Necrosis is always an acute complication, whereas inflammatory complications can occur many years after injection. The rarity of the complications was highlighted, but it was difficult to estimate the incidence and prevalence in the entire population. In addition, concern was raised about serious complications, which, in addition to being permanent, are often untreatable.

In 2009, another series of 18 cases with various complications related to PMMA use highlighted the indiscriminate use of this substance owing to its low cost and the lack of regulation of its sale to nonspecialist physicians and nonphysicians. In 2012,
In the same year, more than 17,000 complications associated with use of PMMA were recorded in Brazil. Meanwhile, the use of nonsurgical procedures (primarily for filling) increased by 390% in a 2-year period.16

**DISCUSSION**

Based on the data for the last decade, concerns about these procedures are inevitable. Despite the significant and immediate results in aesthetic filling, the unregulated popularization of PMMA use, in addition to its use in large volumes with inadequate technique, have shown that PMMA can be harmful when misused for this purpose.

In Brazil, the regulation and supervision of aesthetic medical centers by responsible agencies remain inadequate. In addition, despite the efforts of medical societies, misinformation is widely disseminated, made worse by increasing exposure on social media. This misinformation exposes patients to unsafe procedures.

With reports of significant adverse outcomes in the Brazilian media in recent years, including mutilating and fatal cases, medical societies have spoken out against the use of PMMA for aesthetic purposes. In 2018, both the SBCP and the Brazilian Society of Dermatology (Sociedade Brasileira de Dermatologia, SBD) issued a warning on the use of PMMA, with a contraindication for use in large amounts, reinforcing the unpredictability of results and requesting the recategorization and restriction of its use by ANVISA.17

Reports of adverse outcomes related to PMMA are rare in the medical literature, with complication rates ranging from 0.01% to 3%.2-18 Although small, these numbers deserve attention, since underreporting of complications is known to occur; both because they may be delayed and also due to omission of reporting in the medical record.3 Furthermore, it may be unclear whether complications are caused by PMMA itself or poor technique. Complications caused by permanent fillers deserve special attention, because they create chronic problems and are difficult to treat. Despite the availability of protocols, there is no consensus on standardization of treatment.19

Complications, such as necrosis, are even rarer (0.003%).18 Technical failure is attributed to the application of needles in flat surfaces and not necessarily to PMMA. In one author’s account of personal experience with more than 5,000 cases published in 2012,20 a complication rate of 0.01% was reported, with no necrosis observed. This was ascribed to a histopathological study of 63 cases of complications attributed to PMMA identified 5 cases with acute complications, all of which developed necrosis after injection.18

In 2010, the Regional Medical Council of Paraná (Conselho Regional de Medicina do Paraná, CRM-PR) issued an opinion that the unrestricted use of PMMA in gluteal augmentation, or use in large quantities, was unsafe and unpredictable, and could lead to chronic reactions and unmanageable complications. In 2012, ANVISA issued a safety alert highlighting the possible chronic complications of PMMA, as well as the need for professional training in its use.12,13

In 2013, the CFM issued a new opinion reinforcing the 2010 opinion of the CRM-PR and reaffirming the limited indications for PMMA injection, noting that use in large quantities could lead to unpredictable results. In this opinion, both the Brazilian Society of Plastic Surgery (Sociedade Brasileira de Cirurgia Plástica, SBCP) and the Technical Chamber of Plastic Surgery of the CFM recommended that PMMA only be used by physicians, as well as in small doses and with restrictions.

The 2016 census by the SBCP-SP (regional São Paulo branch) reported that a total of 4,432 procedures were performed to treat complications of PMMA injection, equivalent to 0.7% of the total number of reparative procedures in that year. In the same year, the use of nonsurgical procedures (primarily for filling) increased by 390% in a 2-year period.16

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**Figure 6.** Wound appearance after 3 weeks with homologous grafting and subsequent immediate autologous grafting. a: left lateral; b: posterior; c: right lateral; VIII: after ninth debridement and 3 weeks of homologous grafting over the gluteus maximus; IXa: 1:1.5 mesh graft over gluteus maximus after debridement of hypertrophic granulation and epidermal remnants of previous homologous graft; IXc: appearance of wound bed after debridement of previous homologous graft.

**Figure 7.** A: Sixth postoperative day. Appearance of partial, autologous 1:1.5 skin mesh graft. B: 1-month postoperatively, showing integration of graft and wound.
to adherence to 3 principles: deep plane application, use of a microcannula, and use of pure and certified PMMA.

In the case described herein, both the quality of the product and the technique used were questionable. The combination led to severe complications.

Acute local inflammation within the first hours after treatment was an important warning sign that required close monitoring. With progression to necrosis, aggressive surgical debridement combined with negative pressure therapy were imperative for treatment of inflammation and preparation of the wound bed. Debridement surgery is challenging, because healthy tissue is invaded by necrotic tissue, creating a false impression of satisfactory treatment. Coverage of the wound after proper cleaning is also challenging, however, and sequelae and deformities may not be be fully corrected by plastic surgery.

The use of diagnostic tomography in the present case was a predictor of the extent of necrosis, since it revealed densification of affected tissue, even in areas that still appeared clinically healthy. The findings coincided with the debrided areas. There are no comparable studies in the literature.

**CONCLUSION**

Despite low published complication rates in Brazil, an excessive number of repair procedures are needed to correct complications from use of PMMA. The severity of the reported case highlights the need to combat bad practice by untrained practitioners, as well as the need for greater regulation of the commercialization of PMMA. Complications can lead to death and permanent deformity, and treatment is challenging.

**COLLABORATIONS**

**KTK**  
Analysis and/or data interpretation, conception and design study, data curation, final manuscript approval, investigation, methodology, project administration, realization of operations and/or trials, writing - original draft preparation, writing - review & editing.

**MM**  
Realization of operations and/or trials.

**DAM**  
Realization of operations and/or trials, supervision, writing - review & editing.

**AAMJ**  
Supervision.

**RG**  
Supervision, visualization.

**REFERENCES**


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