Topical Management of Wound: A Narrative Review of Cadexomer Iodine Ointment Versus Povidone Iodine Ointment

Shubham Gupta Jr. 1, Sangita Shinde 2, Raju K. Shinde 1

1. General Surgery, Datta Meghe Institute of Medical Sciences (Deemed to be University), Wardha, IND
2. Pharmacology, Datta Meghe Institute of Medical Sciences (Deemed to be University), Wardha, IND

Corresponding author: Shubham Gupta Jr., sg2638@gmail.com

Abstract

Several iodine formulations have been used for wound care for ages, but still, there exist a number of controversial issues regarding their uses in the present era. Many published studies are available for both povidone iodine (PI) and cadexomer iodine (CI) with conflicting outcomes due to different preparations used and different study types. PI has a broad spectrum of activity including antiseptic properties, anti-inflammatory properties, low cytotoxicity, and good tolerability with the absence of associated resistance. CI is an immobilized iodine molecule in a hydrophilic modified-starch polymer bead with the dual property of cleansing the wound by absorbing the exudate and bactericidal effect by sustained release of iodine molecules over the infected wound. The preparations comprising PI and CI improve wound healing and minimize the bacterial infestation or contamination in various chronic wounds, burns, and ulcers. This review narrates the comparison of CI and PI for the management of wounds in the context of biofilm reduction, wound size reduction, and granulation tissue promotion.

Categories: Family/General Practice, Plastic Surgery, General Surgery
Keywords: topical ointments, cadexomer iodine, povidone iodine, wound, ulcer

Introduction And Background

Wound care, irrespective of being acute or chronic, is a serious public health issue that impacts a large number of people with various types of wounds and costs a large number of resources. Wound care has been found to be a considerable burden on patients, healthcare providers, and the healthcare system as a whole. Wound care is a crucial factor to consider while caring for individuals with chronic wounds on a daily basis [1].

Iodine has been used as one of the most effective antiseptics for reducing infection sequelae for over a century, and topical iodine formulations have been employed for wound care. Lugol’s solution is the most basic form of iodine, and it possesses stinging and caustic properties.

Infection management has long been recognized as a necessity in the treatment of wounds. During the last two decades, there have been numerous advances in the science of wound management. Wound management technologies, research, and the formulation of standards of care based on research and clinical data have all contributed to a positive outcome in wound healing. Current literature, on the other hand, supports the use of topical wound treatments for wound care and management [2].

There are several topical products available, each with its own set of benefits and drawbacks. For example, iodine-based preparations tend to release free iodine when they encounter wound exudates, acting as an antiseptic and controlling infection, aiding wound healing [3].

To overcome the drawback, iodine complexes such as povidone iodine (PI) and cadexomer iodine (CI) are used. PI is a trisiodide and polyvinylpyrrolidone combination. When the paste is absorbed in wound exudates, triiodide is liberated, maintaining the balance between triiodide and the PI complex.

Iodine molecules are trapped in a hydrophilic modified-starch polymer bead called CI. The beads of polymers in CI are swelled up by the wound exudates and gently discharge integrated iodine, preventing its accumulation and thereby preventing iodine-related complications such as allergic contact dermatitis, systemic acidosis, and so on [4-6]. CI ointment showed positive effects on the increased rate of epithelial regeneration, wound contractility, desloughing, formation of granulation tissue, and neovascularization [7].

Keeping this context, we present this narrative review to compare the outcome of PI ointment and CI ointment for wound management and overcome the limitations of conventionally used PI ointment in terms of desloughing agent, increased epithelial regeneration, promotion of granulation tissue formation, wound contractility, and neovascularization.
Review

Povidone iodine: role in wound management

In PI, iodine forms a combination with the synthetic carrier polymer povidone, which has no microbicidal action. The PI complex releases free iodine into solution in an aqueous media, and a balance is maintained, with more free iodine being liberated from the PI reservoir as the iodine-consuming germicidal activity progresses. Povidone-bound iodine can be released as a bactericidal agent. The formulation, concentration, and temperature-dependent equilibrium of povidone-bound iodine to free iodine serves as protection against the suppression of granulation tissue formation and minimizes safety and acceptability issues related to skin exposure to previous elemental iodine formulations.

Iodine’s microbicidal activity involves the inhibition of critical bacterial cellular mechanisms and structures, as well as the oxidation of nucleotides, fatty/amino acids in bacterial cell membranes, and cytosolic enzymes involved in the respiratory chain, resulting in their denaturant and deactivation. At the molecular level, however, the exact sequence of events remains unclear.

In vitro evidence suggests that iodine not only has broad antibacterial properties but also inhibits inflammation caused by infections and the host response. These anti-inflammatory impacts are likely to be multifaceted and clinically significant.

Spectrum of Activity

PI is a topical antibiotic that acts against bacteria, viruses, fungus, spores, protozoa, and amoebic cysts. In conventional antimicrobial testing, PI is seen to kill a number of bacteria strains found to cause nosocomial infections, including methicillin-resistant Staphylococcus aureus (MRSA) and other antibiotic-resistant strains, within 20-30 seconds of exposure. Comparators like chlorhexidine, on the other hand, entail considerably longer exposure times, and most species retain residual bacteria. Because of these properties of PI ointment, it is mostly recommended for ulcerative wounds, small burns, and traumatic skin loss.

Resistance and Cross-Resistance

Resistance to topical and systemic antibiotics like mupirocin, fusidic acid, and gentamicin is on the rise worldwide, as is the prevalence of hospital and community-acquired illnesses. This is a serious medical problem that appears to be caused in part by the overdose and misuse of antibiotics. Clinical strains provide the majority of evidence for bacterial resistance and cross-resistance to antiseptics such as chlorhexidine, quaternary ammonium salts, silver, and triclosan. There appears to be evidence of antiseptic and antibiotic cross-resistance. Systematic testing has found no resistance to PI yet. Iodine has demonstrated no acquired resistance or cross-resistance in over 150 years of use, unlike other antiseptics (with the exception of an apparent lack of cross-resistance to silver). This lack of resistance is most likely due to iodine’s different mechanisms of action.

Activity Against Biofilms

Biofilm is a complex microbiome structure that adheres to the surface and contains different bacterial colonies or single types of cells in a group. These cells, which are embedded in extracellular polymeric substances, a matrix made up primarily of environmental DNA, proteins, and polysaccharides, demonstrated high antibiotic resistance. Iodine solutions are found to get deactivated in the presence of pus, slough, necrotic tissue, and in presence of organic materials present over the wound. In the presence of antimicrobial therapies, biofilms have been shown to slow wound healing and enhance bacterial survival. PI’s long-term efficacy on wound healing in the presence of biofilms was tested in a recent study. PI’s in vitro efficacy against Staphylococcus epidermidis and Staphylococcus aureus growth, as well as the prevention of staphylococcal biofilm formation at sub-inhibitory concentrations, has been verified in studies. Biofilm clearance was higher in this assay than with polyhexamethylene biguanide (PHMB), octenidine, chlorhexidine, mupirocin, and fusidic acid, for example.

Cadexomer iodine: role in wound management

Cadexomer is a 0.9% w/w iodine-containing hydrophilic starch polymer bead. CI releases free iodine (an antiseptic) when it comes into contact with wound exudates, according to a pharmacodynamic study. The product has a dual action: cleansing of the wound and bactericidal action. It also absorbs fluid (up to 6 milliliters/gram of CI) leading to exudate management of the wound by making cleansing easier. Exudative fluid leads to sustained release of iodine particles leading to a bactericidal effect within the dressing for up to 72 hours. It promotes rapid granulation tissue development and accelerates wound healing with substantially less pain and edema. Moreover, CI maintains a moist environment.
to aid in the healing of chronic skin lesions.

**Spectrum of Activity**

Within six or eight weeks of therapy, significant decreases in wound pathogens were observed with CI ointment intervention compared to standard of care (SOC) in randomized controlled trials (RCTs) in cases of venous leg ulcer patients. Hillström [24] used semi-quantitative approaches to show a substantial decline in Staphylococcus aureus ($P < 0.001$) and an improvement in infection. According to Lindsay et al., in the majority of instances, CI ointment treatment resulted in the removal or reduction of organisms, which was linked to decreased odor and ulcer improvement [25]. Skog et al. [26] reported a significant reduction in streptococci, enterococci, and Enterobacteriaceae such as Proteus and Klebsiella species ($P < 0.0001$ and $P < 0.01$, respectively). Thus, CI ointment is advised mainly for chronic exudative wounds like diabetic foot ulcers, venous ulcers, and pressure ulcers where slough, infection, or the risk of infection is a concern [15].

**Activity Against Biofilm**

Biofilms have recently been identified as a source of chronic wound infections and a delay in wound healing, with biofilms found in over 78% of chronic non-healing wounds like venous leg ulcers, diabetic foot ulcers, and pressure ulcers [27]. Biofilm provides a substantial clinical challenge, especially because of its increased antimicrobial tolerance and capacity to elude the human immune response. In vitro models combining several therapeutically relevant circumstances and substrates, such as porcine tissue, have shown that CI ointment is effective against mature biofilms; in addition, animal models indicate encouraging outcomes with CI ointment against biofilm in a wound [28].

**Wound Area Reduction and Wound Healing**

Following CI ointment management, 10 RCTs in the published literature showed significant improvements in wound area reduction or full healing events in chronic wounds. Hillström [24] observed a substantial reduction in wound area with CI ointment after only one week of treatment, which lasted until the completion of the research (six weeks). Furthermore, Skog et al. [26] found that the significant reduction in ulcer size by one and two weeks ($P = 0.005$) translated to a 34% reduction in ulcer size by six weeks, compared to a 5% increase in ulcer size in the SOC group.

**Critical evaluation**

While PI is most commonly recommended for ulcerative wounds, small burns, and traumatic skin loss, CI is administered in chronic exudative wounds like diabetic foot ulcers, venous ulcers, and pressure ulcers where slough, infection, or the risk of infection is a concern, according to multiple studies [15]. The findings of various in vivo investigations show unequivocally that there is sufficient evidence to recommend the use of CI in chronic wounds. According to a review, there is insufficient evidence to demonstrate that CI has a deleterious effect on wound healing and infection [21-27].

PI, on the other hand, showed to be efficient in reducing bacteria counts and preventing wound infections when used in the presence of infection in acute wounds like traumatic lacerations and burn wounds [29-31]. PI inhibits collagen formation, is toxic to fibroblasts and keratinocytes, and inhibits epithelial cell migration, thereby impairing the healing process in non-infected human wounds [9,32-34].

In human clinical studies, CI was found to be the only drug that reduced total microbial burden, including biofilm, to a satisfactory and effective level along with accelerating epithelisation and granulation tissue formation [25,35].

**Reactive Equivalence of Iodine Present in PI Sugar Ointment and CI Ointment**

The optimal iodine concentration for antibacterial activity was 0.01 w/v %, indicating that iodine–L-tyrosine reactivity is closely linked to antimicrobial action. The concentration of free iodine is unrelated to the concentration of total iodine, demonstrating that free but not total iodine is required for antibacterial activity. It is recommended that the concentration of iodine be increased to 0.1 w/v % because a large quantity of iodine is absorbed when used topically at low concentrations (0.01 w/v %) in wounds.

PI sugar ointment interacted more efficiently with L-tyrosine at this therapeutically relevant dose as compared to lecithin, whereas CI ointment reacted more efficiently with lecithin as compared to L-tyrosine. The quantity of iodine in PI sugar ointment that reacted with actual wound exudates was two times higher than the amount of iodine in CI ointment that reacted with wound exudates, alluding that iodine is quickly consumed by the protein component in PI sugar ointment and the antiseptic effect is quickly diminished. Iodine in CI ointment maybe eventually absorbed by protein components, allowing the antimicrobial properties to last longer [36].
**Equivalence of Water Absorption Among PI Sugar Ointment and CI Ointment**

Using an agarose gel to measure water absorption capacity, it was discovered that CI ointment had a 2.9-fold higher water absorption capacity per weight over 24 hours than PI sugar ointment. These findings imply that the amount of water absorbed is affected by whether the base ointment stays in its original form or dissolves after water absorption. When PI sugar ointment and CI ointment are utilized to treat pressure ulcers, PI sugar ointment may have transitory low water absorption since the base ointment gets dissolved, whereas CI ointment may have persistent water absorption as the rate-determining step is the diffusion of dissolved macrogol and water to macromolecular beads [37].

**Conclusions**

A combination of PI and CI enhances healing and reduces bacterial contamination in a variety of chronic wounds, burns, and ulcers. Despite the antibacterial benefits acquired through its use, various possible downsides have been reported in its therapeutic application, with varying and contradictory findings, prompting practitioners to be cautious about using forms of iodine for topical wound treatment. Furthermore, the ability of the base ointments used in PI sugar ointment and CI ointment to absorb water differs. As a result, when these two ointments are administered for pathogenetically similar conditions, they may provide different effects. The literature has revealed that CI is more effective in treating chronic wounds, especially with high levels of exudates as compared to PI. Several studies have strongly supported that CI has a better outcome than PI with respect to desloughing action of pus and debris, reduction of wound size, and promotion of granulation tissue formation leading to better wound healing and management.

**Additional Information**

**Disclosures**

**Conflicts of interest:** In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that no financial support was received from any organization for the submitted work. Financial relationships: All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. Other relationships: All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

**References**

1. Situm M, Kolić M, Redžepi G, Antolić S: Chronic wounds as a public health problem. (Article in Croatian) . Acta Med Croatia. 2014, 68:5-7.
2. Lipsky BA, Hoye C: Topical antimicrobial therapy for treating chronic wounds . Clin Infect Dis. 2009, 49:1541-9. 10.1086/644752
3. Block SS: Disinfection, Sterilization, and Preservation. Lea & Febiger, Philadelphia, PA; 1991.
4. Reyazulla MA, Gopinath AL, Vaibhav N, Raut RP: An unusual complication of late onset allergic contact dermatitis to povidone iodine in oral & maxillofacial surgery – a report of 2 cases. Eur Ann Allergy Clin Immunol. 2014, 46:157-9.
5. Yavancan O, Kara OD, Soren G, Aksu N: Allergic dermatitis caused by povidone iodine: an uncommon complication of chronic peritoneal dialysis treatment. Adv Perit Dial. 2005, 21:131-3.
6. Pietsch J, Meakins JL: Complications of povidone-iodine absorption in topically treated burn patients . Lancet. 1976, 1:280-2. 10.1016/s0140-6736(76)91406-9
7. Lamme EN, Gustafsson T, Middelhoek E: Cadexomer-iodine ointment shows stimulation of epidermal regeneration in experimental full-thickness wounds. Arch Dermatol Res. 1998, 290:18-24. 10.1007/s004030050271
8. Fleischer W, Reimer K: Povidone-iodine in antisepsis - state of the art . Dermatology. 1997, 195:3-9. 10.1159/000246022
9. Rackur H: New aspects of mechanism of action of povidone-iodine . J Hosp Infect. 1985, 6:13-23. 10.1016/s0195-6701(85)80041-4
10. Ripa S, Bruno N, Reder RF, Casillus R, Roth RI: Clinical applications of povidone-iodine as a topical antimicrobial. Handbook of Topical Antimicrobials. CRC Press, Boca Raton, FL; 2002. 22.
11. Kanagalingam J, Feliciano R, Hah HJ, Labib H, Le TA, Lin JC: Practical use of povidone-iodine antiseptic in the maintenance of oral health and in the prevention and treatment of common oropharyngeal infections. Int J Clin Pract. 2015, 69:1247-56. 10.1111/i tcp.12707
12. Al-Kaisy AA, Salih Sahib A: Role of the antioxidant effect of vitamin e with vitamin C and topical povidone-iodine ointment in the treatment of burns. Ann Burns Fire Disasters. 2005, 18:19-30.
13. Cooper RA: Iodine revisited. Int Wound J. 2007, 4:124-37. 10.1111/j.1742-481X.2007.00314.x
14. Yasuda T, Yoshimura Y, Takada H, et al.: Comparison of bactericidal effects of commonly used antiseptics against pathogens causing nosocomial infections. Part 2. Dermatology. 1997, 195:39-28. 10.1159/000246026
15. Khan MN, Naqvi AH: Antiseptics, iodine, povidone iodine and traumatic wound cleansing . J Tissue Viability. 2006, 16:6-10. 10.1016/s0965-206x(06)64002-3
16. Lachapelle M, Castel O, Casado AF, Leroy B, Micali G, Tennystedt D, Lambert J: Antiseptics in the era of bacterial resistance: a focus on povidone iodine. Clin Pract. 2013, 10:579-92.
17. Lanker Klossner B, Widmer HR, Frey F: Nondevelopment of resistance by bacteria during hospital use of povidone-iodine. Dermatology, 1997, 195:10-5. 10.1159/000246024
18. Sharma D, Misba L, Khan AU: Antibiotics versus biofilm: an emerging battleground in microbial communities. Antimicrob Resist Infect Control. 2019, 8:76. 10.1186/s13756-019-0533-3
19. Morgan D: Is there still a role for antiseptics? J Tissue Viability. 1995, 3:30-4. 10.1016/0906-268X(95)80074-0
20. Percival SL, Finnegan S, Donelli G, Vuotto C, Rimmer S, Lipsky BA: Antiseptics for treating infected wounds: efficacy on biofilms and effect of pH. Crit Rev Microbiol. 2016, 42:293-309. 10.3109/1040841X.2014.940495
21. Fitzgerald DJ, Renick PJ, Forrest EC, et al.: Cadexomer iodine provides superior efficacy against bacterial wound biofilms in vitro and in vivo. Wound Repair Regen. 2017, 25:13-24. 10.1111/wrr.12497
22. Roche ED, Woodmansey EJ, Yang Q, Gibson DJ, Zhang H, Schultz GS: Cadexomer iodine effectively reduces bacterial biofilm in porcine wounds ex vivo and in vivo. Int Wound J. 2019, 16:647-85. 10.1111/iwj.13080
23. Holloway GA Jr, Johansen KH, Barnes RW, Pierce GE: Multicenter trial of cadexomer iodine to treat venous stasis ulcer. West J Med. 1989, 151:55-8.
24. Hillström L: Iodosorb compared to standard treatment in chronic venous leg ulcers—a multicenter study. Acta Chir Scand Suppl. 1988, 544:53-6.
25. Lindsay G, Latta D, Lyons KGB, Livingstone ED, Thomson W: A study in general practice of the efficacy of cadexomer iodine in venous leg ulcers treated on alternate days. Acta Ther. 1986, 12:141-8.
26. Skog E, Arnesjö B, Troëng T, et al.: A randomized trial comparing cadexomer iodine and standard treatment in the out-patient management of chronic venous ulcers. Br J Dermatol. 1983, 109:77-85. 10.1111/j.1365-2133.1983.tb03995.x
27. Woo K, Dowsett C, Costa B, Ebohon S, Woodmansey EJ, Malone M: Efficacy of topical cadexomer iodine treatment in chronic wounds: systematic review and meta-analysis of comparative clinical trials. Int Wound J. 2021, 18:586-97. 10.1111/iwj.13560
28. Stewart PS, Costerton JW: Antibiotic resistance of bacteria in biofilms. Lancet. 2001, 358:135-8. 10.1016/s0140-6736(01)05321-1
29. Gravett A, Sterner S, Clinton JE, Ruiz E: A trial of povidone-iodine in the prevention of infection in sutured lacerations. Ann Emerg Med. 1987, 16:167-71. 10.1016/s0196-0644(87)80008-2
30. Goldenheim PD: An appraisal of povidone-iodine and wound healing. Postgrad Med J. 1993, 69:97-105.
31. Gordon J: Clinical significance of methicillin-sensitive and methicillin-resistant Staphylococcus aureus in UK hospitals and the relevance of povidone-iodine in their control. Postgrad Med J. 1993, 69:910-16.
32. Van den Broek PJ, Buys LF, Van Furth R: Interaction of povidone-iodine compounds, phagocytic cells, and microorganisms. Antimicrob Agents Chemother. 1982, 22:595-7. 10.1128/AAC.22.4.595
33. Connolly JC, Gilmore OJ: A study of the effect of povidone-iodine on polymorphonuclear leucocyte chemotaxis. Br J Exp Pathol. 1979, 60:662-6.
34. Ninnemann JL, Stein MD: Suppressor cell induction by povidone-iodine: in vitro demonstration of a consequence of clinical burn treatment with betadine. J Immunol. 1981, 126:1905-8.
35. Mertz P, Davis S, Brewer L, Franzen L: Can antimicrobials be effective without impairing wound healing? The evaluation of a cadexomer iodine ointment. Wounds. 1994, 6:184-93.
36. Berkelman RL, Holland BW, Anderson RL: Increased bactericidal activity of dilute preparations of povidone-iodine solutions. J Clin Microbiol. 1982, 15:655-9. 10.1128/jcm.15.5.655-659.1982
37. Zhou LH, Nahm WK, Badiavas E, Yufit T, Falanga V: Slow release iodine preparation and wound healing: in vitro effects consistent with lack of in vivo toxicity in human chronic wounds. Br J Dermatol. 2002, 146:565-74. 10.1046/j.1365-2133.2002.04605.x