Zinc pyrithione in alcohol-based products for skin antisepsis: Persistence of antimicrobial effects

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Alcohol-based products for skin antisepsis have enjoyed a long history of safety and efficacy in the United States and abroad. Examples of use include surgical scrubs, health care personnel handwashes, patient preoperative skin preparations, injection/catheter site preparations, preoperative antiseptic shower solutions, hand rubs/sanitizers, and tinctures such as those of iodine. Of the antimicrobials routinely used for skin antisepsis (alcohols, chlorhexidine gluconate [CHG], iodine/iodophors, parachlorometaxylenol [PCMX], triclosan, and quaternary ammonium compounds), alcohols are by far the fastest acting and most efficacious. Almost exclusively, the short-chain, aliphatic alcohols—ethanol, isopropanol, and, in Europe, n-propanol—are used for skin and hand antisepsis. They have excellent activity against bacteria, fungi, and enveloped (and some nonenveloped) viruses. Alcohols may be used either alone or in combination with other antimicrobials to increase the efficacy and confer substantivity (persistence).

The Centers for Disease Control (CDC) and Prevention in its “Guideline for Prevention of Surgical Site Infection, 1999” acknowledged the utility of alcohol-based products in preoperative hand/forearm antisepsis and preoperative skin preparation. In late 2002, the recommendations of the Healthcare Infection Control Practices Advisory Committee and the HICPAC/SHEA/APIC/IDSA Hand Hygiene Task Force were published. This report recognized the speed and broad antimicrobial spectrum of alcohol, codified uses for alcohol-based products in skin antisepsis practices, and noted the lack of antimicrobial persistence associated with alcohol when used alone. The FDA also noted alcohol’s lack of antimicrobial persistence; however, it allows a preservative agent to be incorporated into the vehicle (defined as the product without the active ingredient) to provide for the persistent antimicrobial effect necessary to sustain a reduction in the number of bacteria for 6 hours postapplication. The preservative supplies the required persistent antimicrobial activity for these formulations to meet or exceed the established agency efficacy. Preservative systems in some of the currently marketed alcohol-based products for skin antisepsis include iodine/iodophors compounds, CHG, and zinc pyrithione (ZPT). Although much has been written about the antimicrobial effects and the preservative potential of iodine and CHG, little has been published regarding these characteristics as attributes of ZPT. Therefore, this article includes a brief review of the published literature related to the development, safety,
efficacy, and clinical utility of ZPT and evaluating its merits as a preservative for alcohol-based products of skin antisepsis.

CHARACTERISTICS OF ZINC PYRITHIONE

Synthesis, structure, and solubility

The natural antibiotic aspergillic acid contains a cyclic hydroxamic acid functional group in a pyrazine nucleus. Attempts to develop synthetic methods for introducing heterocyclic rings into the hydroxamic acid group present in aspergillic acid led in 1950 to the preparation for N-hydroxy-2-pyridinethione (HPT). The synthesis was achieved by conversion of a 2-pyridyl ether to its N-oxide, followed by dealkylation.13 Reaction of 2-bromopyridine-N-oxide with thiourea forms 2-pyridyl-N-oxide-isothiourea hydrobromide and, followed by treatment with aqueous sodium carbonate, produces N-hydroxy-2-pyridinethione (HPT).14 This compound was shown to have potent antimicrobial properties. In vitro, 1 μg HPT would inhibit Staphylococcus aureus.15 Thus, this synthetic analog was 30 times more potent as an antimicrobial than the native aspergillic acid. Later, HPT was shown to have extremely potent activity against gram-positive and gram-negative bacterial species as well as strong activity against yeasts and fungi: e.g., Aspergillus, Trichophyton species, Candida albicans, and Cryptococcus species.16 Cox, in the mid-1950s, reviewed the uses of pyridine-N-oxides and noted that the mercapto derivatives formed quaternary ammonium compounds or heavy metal salts, which were described as effective fungicides and antibacterial agents and suggested their uses in pharmaceutic preparations.17 As a zinc chelate of HPT, ZPT exists in the monomeric form as 2 pyridine rings bound to a central zinc atom by bonds between the zinc atom and the sulphur and oxygen molecules of the pyridine ring structures.18,19 ZPT is practically insoluble in water, organic solvents, or surfactants—a property that can be problematical when formulating with the compound.20 The structure of ZPT is presented in Fig 1.

Applications

ZPT has a long history of numerous medical, scientific, and industrial applications. The compound has a master file registered with the FDA. Of some 1350 compounds screened, ZPT was one of the most active antifungal and antibacterial compounds examined.20,21 For example, Pityrosporum ovale is thought to play a pathogenic role in skin conditions such as seborrheic dermatitis and dandruff.22 This led one well-known pharmaceutic company to conduct safety evaluations of ZPT in a shampoo formulation.21 The result was an antidandruff product, Head & Shoulders (Procter & Gamble, Cincinnati, Ohio) that is currently sold over-the-counter with 1.0% ZPT as the active ingredient. Presently, the FDA acknowledges the efficacy of ZPT in the treatment of dandruff and seborrheic dermatitis.23,24 The ZPT-based shampoo has shown to be effective in treating tinea versicolor,25 and it is highly effective for psoriasis of the scalp.26 Cosmetic preservation has been a frequent use of ZPT for many years.18 ZPT has proven to be effective, even at low concentrations, against both gram-positive and gram-negative bacteria and fungi. It is compatible with most commonly used cosmetic ingredients; has a good toxicity profile for this type of application; and can be safely used to minimize discoloration, off-odors, and emulsion breaks because of microorganisms. ZPT helps to prevent spoilage of cosmetics because of microbial contamination during use, is found in GoJo Hand Cleaner products (GOJO Industries, Inc. Akron, Ohio), and in over-the-counter antimicrobials such as Lanacane (Combe, Inc., White Plains, NY).27 Finally, ZPT has also been used in commercial laundries for inhibiting mold growth on fabrics.20

Safety and toxicity

In addition to a 50-year product history, the safety and toxicity of ZPT has been formally evaluated in
animal models and by in vitro and human in vivo studies. When ZPT was applied to intact skin of monkeys with surfactants, the absorption was only 0.20% of the amount applied.\textsuperscript{28} Concentrations of ZPT in the blood stream following topical application are below the threshold of detection. The absorbed dose would likely be further reduced in humans by the fact that hand skin has a lower permeability than scalp skin.\textsuperscript{29,30} ZPT does not induce primary skin irritation or sensitization in human skin.\textsuperscript{30} Finally, when a 2.0% suspension of ZPT was instilled into the conjunctival sacs of the eyes of 6 albino rabbits, ZPT produced only slight irritation of the conjunctival vessels, which lasted 3 days, but no corneal opacities were observed.\textsuperscript{31} Ocular toxicity is well known to be associated with other preservative systems such as CHG, and skin irritation and sensitization with iodine and CHG-based products has been documented.\textsuperscript{10,32-34}

### Antimicrobial activity

The available data on the mode of antimicrobial action suggest that ZPT is membrane active, as indicated by the inhibition of uptake of several unrelated substrates in both bacteria and fungi\textsuperscript{35-38} and the observed depolarization of the transmembrane potential in \textit{Neurospora crassa}.\textsuperscript{39} The effects of an antimicrobial agent on substrate transport and related metabolism may be used as indicators of the membrane activity of the test agent.\textsuperscript{40} In turn, these effects may be reflected as a reduction in intracellular ATP levels.\textsuperscript{41} ZPT is a poor inhibitor of substrate catabolism. Subinhibitory concentrations of the biocide greatly reduce intracellular ATP levels in both \textit{Escherichia coli} and \textit{Pseudomonas aeruginosa}. This is thought to be due to the action of ZPT on the gram-negative bacterial membrane.\textsuperscript{42} Further investigation of the action at the membrane suggests that ZPT forms stable interactions with the bacterial membrane phospholipid phosphatidylethanolamine. This may result in the disaggregation of the phospholipid head structure at the outer membrane and may also indicate chelation of phosphatidylethanolamine head groups from the core structure of the external lipopolysaccharide. This would further disrupt the membrane.\textsuperscript{43} In addition, current-voltage analysis demonstrates that the depolarization of the bacterial membrane is accompanied by a decrease in membrane electrical conductance in a manner consistent with inhibition of the primary proton pump and consistent with a mode of action of ZPT on plasma membrane ion channels. Therefore, ZPT inhibits membrane transport via a direct or indirect effect on the primary proton pump that energizes transport, and the site of action of ZPT is likely to be intracellular rather than extracellular.\textsuperscript{42} Other studies on the mode of action of pyridine-N-oxides has demonstrated their potent bactericidal activity to be linked to their ability to chelate; i.e., to form cyclic complexes with the ions of heavy metals.\textsuperscript{44} Additional investigations reported by Hyde and Nelson have suggested that other mechanisms may be applicable.\textsuperscript{18} The authors propose that the pyrithione is an antimetabolite of the pyridine derivative pyridoxal and suggest that the activity of ZPT may be analogous to the inhibition of microbial folate production by sulfa drugs. Finally, dipole structure of the molecule creates a pseudoquaternary ammonium group, providing yet another potential mode of antimicrobial action for ZPT. Multiple mechanisms appear to be at work, suggesting that antimicrobial resistance is unlikely to develop. Reports of the development of antimicrobial resistance are not readily available, and further detailed investigation may be appropriate.

**In vitro efficacy data.** The in vitro composite data in Table 1 document the broad spectrum of antimicrobial activity of ZPT as well as quantify the inhibitory capacity of this compound when tested against numerous bacteria, yeast, and fungi (Personal communications from Ron Jones, M. D., Professor and Director, Division of Medical Microbiology, Director, Anti-Infectives Research Center and Special Microbiology Laboratories, Department of Pathology, University of Iowa College of Medicine, Iowa City, Ia, January 1993).\textsuperscript{20,22,27,42} The antimicrobial range of ZPT is sufficient to include many pathogenic, opportunistic, and saprophytic organisms. Inhibitory concentrations are often achieved at a level of 40 μg/mL (40 ppm) or less for numerous pathogenic species.

**In vivo efficacy and persistence data.** In 1979, Leyden et al\textsuperscript{45} developed a novel in vivo assay now known as the persistence (substantivity) test. This significant development was used to demonstrate the in vivo efficacy of ZPT in topical antisepsis applications. The assay determines the ability of the test agent to establish a reservoir in the stratum corneum. Substantive agents that diffuse into the stratum corneum or bind chemically to it will not be readily removed either by loss from the surface or by absorption. The antibacterial effect will therefore last several days. For this assay, 0.5 mL of the test agent is applied with a pipette twice daily for 4 consecutive days to the entire volar forearm. Twenty-four hours after the last application, occlusive dressings are applied. If an antimicrobial effect is demonstrable, the test is repeated, and the posttreatment interval is extended to 72 hours. The geometric means for bacterial counts per square centimeter are determined. For this study, 1.0% ZPT was compared with an equal concentration of CHG. The results of this 8-subject study demonstrated antimicrobial parity at 24 hours; however, at the
Table 1. Mean minimal inhibitory concentration of ZPT for selected bacteria, yeast, and fungi

<table>
<thead>
<tr>
<th>Organism tested</th>
<th>Number of strains/strain identification</th>
<th>MIC 50 in ppm</th>
<th>MIC 90 in ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Staphylococcus aureus</td>
<td>10 strains</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>CNS</td>
<td>10 strains</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Enterococcus species</td>
<td>10 strains</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>Streptococcus faecalis</td>
<td>ATCC 19433</td>
<td>10</td>
<td>20</td>
</tr>
<tr>
<td>Bacillus cereus</td>
<td>ATCC 11778</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Sarcina lutea</td>
<td>ATCC 9341</td>
<td>10</td>
<td>20</td>
</tr>
<tr>
<td>Escherichia coli</td>
<td>10 strains</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Proteus vulgaris</td>
<td>ATCC 9920</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Pseudomonas aeruginosa</td>
<td>NCIMB 10548</td>
<td>ND</td>
<td>13</td>
</tr>
<tr>
<td>Citrobacter species</td>
<td>10 strains</td>
<td>20</td>
<td>40</td>
</tr>
<tr>
<td>Acinetobacter species</td>
<td>10 strains</td>
<td>40</td>
<td>150</td>
</tr>
<tr>
<td>Serratia marcescens</td>
<td>10 strains</td>
<td>20</td>
<td>40</td>
</tr>
<tr>
<td>Salmonella/Shigella</td>
<td>10 strains</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>Xanthomonas maltophilia</td>
<td>10 strains</td>
<td>150</td>
<td>300</td>
</tr>
<tr>
<td>Morganella morganii</td>
<td>5 strains</td>
<td>40</td>
<td>150</td>
</tr>
<tr>
<td>Enterobacter species</td>
<td>10 strains</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>Klebsiella species</td>
<td>10 strains</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>Dermatophytes</td>
<td>5 strains</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Aspergillus species</td>
<td>5 strains</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Candida species</td>
<td>5 strains</td>
<td>10</td>
<td>10</td>
</tr>
</tbody>
</table>

MIC. Mean minimal inhibitory concentration; ND, not done. ppm, parts per million.

Includes S epidermidis (4 strains), S homina (1 strain), S saprophyticus (3 strains), and S simulans (2 strains).

Includes C difficile (5 strains), E fecalis (2 strains), E faecium (2 strains), E avium (1 strain), E durans (1 strain), and E raffinosus (1 strain).

Includes C diversus (6 strains) and C freundi (4 strains).

Includes A oritatus (8 strains) and A lewffi (2 strains).

Includes S enteticus (6 strains) and S sonnei (4 strains).

Includes E cloacae (7 strains) and E aerogenes (3 strains).

Includes K oxytoca (2 strains) and K pneumoniae (8 strains).

Includes I strain each of M canis, M gypseum, T mentagrophytes, T rubrum, and Tichyphilopont spp.

Includes A flavus (2 strains), A fumigatus (2 strains), and A terreus (1 strain).

Includes C glabrata (1 strain), C krusei (1 strain), C lusitaniae (1 strain), C parapsilosis (1 strain), and C tropicalis (1 strain).

Table 2. Expanded flora test: bacterial counts/cm² forearm skin for 1.0% ZPT versus 70% ethanol

<table>
<thead>
<tr>
<th>Antimicrobial</th>
<th>Sample time postmicrobial expansion</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>6 hours</td>
</tr>
<tr>
<td>ZPT 1.0%</td>
<td>65,110</td>
</tr>
<tr>
<td>Ethanol 70%</td>
<td>14,000</td>
</tr>
<tr>
<td>Control untreated</td>
<td>1,320,000</td>
</tr>
</tbody>
</table>

MIC, not done. ppm, parts per million.

Table 2 data obtained at the 24- and 48-hour sample points.

Persistence mechanism

Following topical application, ZPT is deposited in the epidermis; however, the largest proportion of ZPT is deposited on the outer layers of the stratum corneum. ZPT is soluble in sebum, and, when applied to the skin, the ZPT in sebum is localized in the hair follicles. No transepidermal penetration occurs. The articles provided do not suggest whether the “dissolution” is of the intact ZPT or whether the zinc is first chelated by a separate mechanism, thus leaving the more soluble pyrithione available to penetrate via the follicle. Others used ZPT with radioactive sulfur to monitor tissue location so that the pyrithione moiety would be observed. Deposited particles of ZPT appear to adhere firmly to the hair and stratum corneum and to withstand copious rinsing. These adherent particles of ZPT are not removed by a standard wash and rinse procedure.
particles probably act as a reservoir and are responsible for the prolonged antimicrobial effect (persistence). Diminution of the amount of antibacterial effect with time is probably due to a great extent to normal turnover of the stratum corneum. Also, solubilized ZPT would eventually purge and/or break down further. It is pertinent to note that sebum and topical lipids can inactivate many antimicrobials; examples include hexachlorophene and benzethonium chloride. ZPT is not only soluble in sebum but retains its antimicrobial activity.

**Zinc pyrithione formulations with alcohol**

Given the significant history, safety, and diversity of ZPT, it is only logical that other significant uses would be found for this antimicrobial. Despite the limited solubility and the formulation challenges that are characteristic of the compound, ZPT has been used successfully and most recently as a preservative system for several marketed alcohol-based products of skin antisepsis, at concentrations of 0.25% for waterless applications and up to 1.0% for water-aided products. These include surgical scrubs, health care personnel handwashes, patient preoperative skin preparations, and preoperative bath or shower solutions. In each case, these products are compliant with the FDA’s OTC Drug Monograph and command in vitro and in vivo antimicrobial efficacy as required for immediacy of kill, persistence, and residual effect.

**Hobson et al.** was the first to document in vivo the value of ZPT as a preservative system for an alcohol-based, water-aided surgical scrub formulation. The study compared the efficacy of 70% alcohol preserved with ZPT to a second scrub containing only 4.0% CHG and a third scrub containing only 7.5% iodine. Their data were obtained following 11 scrubs over a 5-day period. The immediacy of kill provided by the alcohol-based product on day 1 was readily demonstrated by the nearly 3.0-log10 reduction from baseline counts. The antimicrobial effect becomes enhanced on days 2 and 5 as a result of the contributions provided in part by the preservative system. At the end of the 5-day test period, the superiority of alcohol preserved with ZPT was readily apparent (4.78-log10 reduction from baseline) when compared with CHG or iodine-based products (3.43- and 1.27-log10 reduction from baseline, respectively) for the same day 5 time point. In a second in vivo study, a different formulation of an alcohol-based, ZPT-preserved, waterless surgical scrub was compared with another waterless, alcohol-based scrub containing 1.0% CHG as a preservative (Avagard, 3M Health Care, St Paul, Minn). As may be observed in **Fig 2**, the products performed similarly on days 1 and 2 (alcohol/ZPT log10 reductions from baseline = 2.32 and 3.07, respectively; alcohol/CHG log10 reductions from baseline = 2.08 and 2.87, respectively). However, by day 5, the antimicrobial efficacy of the alcohol/ZPT product was notably better (log10 reduction from baseline = 3.68) than that of the alcohol/CHG product (log10 reduction from baseline = 3.09) because the ZPT-preserved formulation more readily exceeded the 3.0-log10 reduction required for the surgical scrub indication. In a separate in vivo study, the cosmetic and skin-conditioning properties of the 2 waterless formulations were judged to be equal (personal communication, Dr. Ronald Rizer; manuscript submitted for publication to AORN Journal).

Additional testing was performed with the waterless alcohol/ZPT formulation to assess its virucidal capacity against some common human pathogens. These included human coronavirus (HCoV, ATCC VR-740, both HCoV and SARS-HCoV belong to the virus family Coronaviridae), the human immunodeficiency virus type 1 (HIV-1, from Zepto Metrix Corp. of Buffalo, NY), hepatitis A virus (HAV, from CREM, University of Ottawa), herpes simplex type 1 (HSV-1, ATCC VR-260), and human rotavirus (strain Wa, ATCC VR-2018). These agents represent both RNA and DNA viruses, some with envelopes and others without. The HAV and the human rotavirus represent 2 nonenveloped RNA agents (one
linear single-stranded, the other segmented double stranded) that are relatively resistant to known antimicrobials. Using standard virologic techniques, the viral agents were cultured in cell lines appropriate for propagation (see Table 3) to levels at or above the required titer of $10^6.00$ infectious units/mL. The viruses were then harvested and exposed for 3 minutes (30 seconds for the human coronavirus) to the test formulation diluted by virus inoculum to a 90% concentration. Following exposure to the test product, the residual infectious viral population was determined for each virus, employing techniques appropriate for that virus. All testing was performed per American Society for Testing and Materials (ASTM) method E 1052-96, a Standard Test Method for Efficacy of Antimicrobial Agents Against Viruses in Suspension.50

The results of this effort are summarized in Table 3. The test criterion for classification as a virucide is not less than a 3.0-log$_{10}$ reduction in the postexposure virus population, beyond any observed cytotoxicity. Exposure to the product produced a 3- to 4-log$_{10}$ reduction in all virus populations tested, thereby meeting or exceeding the standard for classification as a virucide.

Seal and Paul-Cheadle confirmed in vivo the antimicrobial efficacy and thus the value of alcohol products formulated with ZPT for added persistence.5 Their report demonstrated the superioriority of an alcohol-based, ZPT-preserved, preoperative shower solution used in combination with a similarly formulated patient preoperative skin preparation. The combination of an alcohol-based, ZPT-preserved, preoperative shower solution and a similarly formulated patient preoperative skin preparation was compared with an iodine-based system (7.5% scrub and 10.0% PVPI paint) using identical application schedules in a well-controlled, human in vivo study. For this study, the subjects refrained from the use of any antimicrobial products for 2 weeks to allow for stabilization of the normal skin flora. Pretreatment cultures were obtained of the selected sites (groin) to obtain “baseline” of skin flora, to which posttreatment data could be compared and log$_{10}$ reduction values calculated. The subjects then were washed with the preoperative shower products at T = −12 and T = −6 hours. The preoperative surgical site preparations were applied at T = 0, and samples were collected at 10 minutes and from 6 hours out to 72 hours postapplication of the surgical site preparation. The alcohol-based products demonstrated superior efficacy at nearly every time point. At 72 hours (3 days) following the last product application, the alcohol-based, ZPT-preserved system continued to provide significant antimicrobial action as documented by a 2.0-log$_{10}$ or a 99.0% reduction in the colony-forming units present at baseline. This compares with a <0.50-log$_{10}$ reduction for the same time point with the iodine-based products. The extended period of antimicrobial persistence that is associated with the ZPT-preserved, alcohol-based system would allow for skin closure in a prolonged state of antisepsis and could result in a lower incidence of surgical site infections.

**DISCUSSION**

Alcohol provides fast-acting, broad-spectrum antimicrobial activity. However, the requirement for persistence in topical antiseptics cannot be achieved with alcohol alone. Chlorhexidine gluconate, quaternary ammonium compounds, iodine, and triclosan have persistence features and have been added to alcohol solutions for antisepsis in Europe and/or the United States for many years. Antimicrobial persistence has been reported with chlorhexidine gluconate and triclosan, but it is slow to develop and is not always profound.32,34,46

As a preservative (persistence agent) for topical antisepsis, ZPT was found to have a safety profile and/or antimicrobial efficacy that exceeds iodine, chlorhexidine gluconate, and triclosan.1,5,32,46,49,50 For these reasons, the antimicrobial persistence associated with ZPT has recently been incorporated successfully into commercially acceptable, FDA-compliant, patented products for surgical hand antisepsis and preoperative antiseptics. In conclusion, ZPT is a safe and effective antimicrobial suitable for use as a preservative system with alcohol formulations. The data confirm

### Table 3. Virucidal effects of an alcohol-based, ZPT-preserved, surgical scrub

<table>
<thead>
<tr>
<th>Virus</th>
<th>Virus classification</th>
<th>Cell line</th>
<th>Initial CCID$_{50}$/mL</th>
<th>Posttreatment log$_{10}$ reduction</th>
<th>Detection method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human coronavirus (SARS virus family)</td>
<td>SS RNA enveloped</td>
<td>MRC-5</td>
<td>$10^6.50$</td>
<td>3.0 Plaque assay</td>
<td></td>
</tr>
<tr>
<td>Herpes simplex virus</td>
<td>DS DNA enveloped</td>
<td>VERO</td>
<td>$10^6.67$</td>
<td>3.17 CPE *</td>
<td>Plaque assay</td>
</tr>
<tr>
<td>Human immunodeficiency virus</td>
<td>SS RNA enveloped</td>
<td>CEM</td>
<td>$10^7.50$</td>
<td>4.0 ELISA for p24 antigen</td>
<td></td>
</tr>
<tr>
<td>Hepatitis A virus</td>
<td>SS RNA nonenveloped</td>
<td>FRhK-4</td>
<td>$10^6.50$</td>
<td>3.0 CPE *</td>
<td></td>
</tr>
<tr>
<td>Human rotavirus</td>
<td>DS RNA nonenveloped</td>
<td>MA-104</td>
<td>$10^6.50$</td>
<td>3.0 Plaque assay</td>
<td></td>
</tr>
</tbody>
</table>

*Cytopathic effects.
that ZPT contributes positively toward the overall antimicrobial efficacy of alcohol-based products in which it is used. Additional studies would prove useful in verifying the clinical relevance of this observation. Finally, it is likely that additional uses for this antimicrobial will be found as we continue the struggle with ever increasing resistance to current antibiotics and antimicrobials.

References


Correction
In the article entitled “Private sector hospital response to the 2003 dengue outbreak in the Indian capital metropolis of Delhi” (Am J Infect Control 2004;32:489-92), fifth author P. Maheshwari’s academic designation was presented incorrectly. The author’s designation should have appeared as follows: P. Maheshwari, MRCP.