Organotin(IV) Complexes with 2-Acetylpyridine Benzoyl Hydrazones: Antimicrobial Activity

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Reações de tricloreto de *n*-butilestanho, [BuⁿSnCl₃], e tricloreto de fenilestanho, [PhSnCl₃], com 2-acetilpiridina benzoil hidrazona (H2AcPh), 2-acetilpiridina-*para*-cloro-benzoil hidrazona (H2Ac*p*ClPh) e 2-acetilpiridina-*para*-nitro benzoil hidrazona (H2Ac*p*NO₂Ph) produziram os complexos [BuⁿSn(2AcPh)Cl₂] (1), [BuⁿSn(2Ac*p*ClPh)Cl₂] (2), [BuⁿSn(2Ac*p*NO₂Ph)Cl₂] (3), [PhSn(2AcPh)Cl₂] (4), [PhSn(2Ac*p*ClPh)Cl₂] (5) e [PhSn(2Ac*p*NO₂Ph)Cl₂] (6). H2Ac*p*ClPh revelou-se a hidrazona mais ativa contra *Staphylococcus aureus* e *Candida albicans*. Pela coordenação a atividade antibacteriana tanto do metal quanto das hidrazonas aumenta significativamente. Os complexos 2 e 5 revelaram-se os mais ativos como agentes antimicrobianos.

Reaction of *n*-butyltin trichloride, [BuⁿSnCl₃], and phenyltin trichloride, [PhSnCl₃], with 2-acetylpyridine benzoyl hydrazone (H2AcPh), 2-acetylpyridine *para*-chloro-benzoyl hydrazone (H2AcpClPh) and 2-acetylpyridine *para*-nitro-benzoyl hydrazone (H2AcpNO₂Ph) gave [BuⁿSn(2AcPh)Cl₂] (1), [BuⁿSn(2AcpClPh)Cl₂] (2), [BuⁿSn(2AcpNO₂Ph)Cl₂] (3), [PhSn(2AcPh)Cl₂] (4), [PhSn(2AcpClPh)Cl₂] (5) and [PhSn(2AcpNO₂Ph)Cl₂] (6) as products. Among the hydrazones H2AcpClPh proved to be the most active against *Staphylococcus aureus* and *Candida albicans*. Upon coordination the antibacterial activity of both tin and the hydrazone significantly increases. Complexes 2 and 5 revealed to be the most active as antimicrobial agents.

Keywords: 2-acetylpyridine benzoyl hydrazones, organotin(IV) complexes, crystal structures, antimicrobial activity

Introduction

Hydrazones are a versatile class of compounds which present a wide range of biological applications as antimicrobial,¹ antitubecular,² anticonvulsant,³ antiinflammatory,⁴ cytotoxic⁵ and vasodilator⁶ agents. Moreover, 9-acridanone-hydrazones proved to be active against *Schistosoma mansoni*⁷ and quinoxaline-*N*acylhydrazones showed trypanocidal activity.⁸ In addition, methyl pyrazinylketone isonicotinoyl hydrazones have been synthesized in an attempt to develop novel chelators with high affinity for iron for the treatment of iron overload disease.⁹

Hydrazones also proved to be useful as sensitive analytical reagents for the determination of trace amounts of metal ions.^{10,11} Metal complexes with hydrazones have

*e-mail: hberaldo@ufmg.br *On leave from Universidad de Oriente, Santiago de Cuba, Cuba potential applications as catalysts,¹² luminescent probes,¹³ and molecular sensors.¹⁴ Moreover, metal complexes with hydrazones present antimicrobial,¹⁵⁻¹⁷ DNA-binding and cytotoxic activities.¹⁷ It has also been shown that metal complexes with hydrazones can be potent inhibitors of cell growth and DNA syntheses.¹⁸

Tin compounds present applications as PVC stabilizers,¹⁹ for chemical vapor decomposition (CVD),²⁰ in non-linear optics²¹ and in catalyses.²² Organotin complexes present innumerous pharmacological applications as antitumorals,²³ antimicrobials^{24,25} and biocides.²⁶ The cytotoxic activity of a variety of organotin complexes against tumor cell lines has been demonstrated by some of us.^{27,28}

In recent works we started an investigation on metal complexes of pyridine-derived hydrazones.²⁹⁻³¹ In the present paper organotin(IV) complexes were prepared by reacting $[Bu^nSnCl_3]$ and $[PhSnCl_3]$ with 2-acetylpyridine-benzoyl hydrazone (H2AcPh), 2-acetylpyridine *para*-chlorobenzoyl hydrazone (H2AcpClPh) and 2-acetylpyridine

para-nitro-benzoyl hydrazone (H2AcpNO₂Ph) (see Figure 1). The antimicrobial activity of the hydrazones and their organotin(IV) complexes was evaluated against *Candida albicans* and *Staphylococcus aureus*.

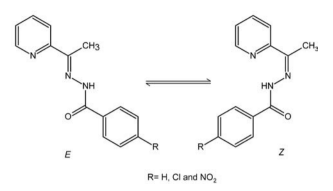


Figure 1. Generic structure of 2-acetylpyridine benzoyl hydrazones (*E* and *Z* configurational isomers).

Experimental

Apparatus

Partial elemental analyses were performed on a Perkin Elmer CHN 2400 analyzer. Infrared spectra were recorded on a Perkin Elmer FT-IR Spectrum GX spectrometer using CsI pellets; an YSI model 31 conductivity bridge was employed for molar conductivity measurements; NMR spectra were obtained with a Bruker DPX-200 (200 MHz) spectrometer or with a Bruker DRX-400 Avance (400 MHz) spectrometer using deuterated dimethylsulfoxide $(dmso-d_{s})$ or deuterated chloroform $(CDCl_{s})$. The ¹H and ¹³C NMR chemical shifts in ppm are reported from internal tetramethylsilane (tms) on the % scale. The ¹¹⁹Sn NMR spectra were measured relative to $Sn(CH_3)_4$. Bruker standard experiments were performed using decoupling sequences for ¹³C and ¹¹⁹Sn. Splitting patterns are designated as follows: s singlet; d doublet; t triplet; q quartet; m multiplet.

n-Butyltin trichloride, phenyltin trichloride, 2-acetylpyridine, benzoyl hidrazide, 4-chloro benzoyl hidrazide, and 4-nitro benzoyl hidrazide were purchased from Sigma-Aldrich and used without further purification. All other chemicals and solvents were of analytical grade.

Syntheses of 2-acetylpyridine-benzoyllhydrazone (H2AcPh), 2-acetylpyridine-para-chloro-benzoyl hydrazone (H2AcpClPh) and 2-acetylpyridine-para-nitro-benzoyl hydrazone (H2AcpNO,Ph)

2-Acetylpyridine-benzoyl hydrazone (H2AcPh) and 2-acetylpyridine-*para*-chloro-benzoyl hydrazone

(H2Ac*p*ClPh) were prepared as described in the literature.^{32,33} 2-Acetylpyridine-*para*-nitro-benzoyl hydrazone (H2Ac*p*NO₂Ph) was obtained by mixing equimolar amounts of 2-acetylpyridine with the *para*-nitro-benzoyl hydrazide in methanol with addition of three drops of acetic acid as catalyst. The reaction mixture was kept under reflux for 6 h. After cooling to room temperature the resulting solid was filtered off, washed with ethanol and ether and dried *in vaccuum*.

*H2AcpNO*₂*Ph*: Color: Beige; Yield: 88%; mp 220-224 °C. Selected IR bands (v_{max} /cm⁻¹): v(NH) 3188 m; v(C=O) 1676 s; v(C=C) + v(CN) 1580 m; 1458 m; 1429 m; ρ(py) 615 m. ¹H NMR (dmso- d_6 , 400 MHz, δ ppm): 11.21 (s, 1H, NH); 8.64 (d, 1H, H6); 8.38 [d, 2H, H(11,13)]; 8.19 [d, 2H, H(10,14)]; 8.16(d,1H, H3); 7.91 (t, 2H, H4); 7.47(t, 1H, H5); 2.50 (s, 3H, CH₃). ¹³C NMR (dmso- d_6 , 400 MHz, ppm): δ 162.69(C6); 154.76 (C7); 149.10 (C2); 148.57(C6); 139.61(C12); 136.61 (C4); 134.06(C9); 129.52(C10,14); 124.25 (C5); 123.30(C11,13); 120.43 (C3).

Synthesis of [*Bu*ⁿ*Sn*(2*AcPh*)*Cl*₂] (1), [*Bu*ⁿ*Sn*(2*AcpClPh*)*Cl*₂] (2), [*Bu*ⁿ*Sn*(2*AcpNO*₂*Ph*)*Cl*₂] (3), [*PhSn*(2*AcPh*)*Cl*₂] (4), [*PhSn*(2*AcpClPh*)*Cl*₂] (5) and [*PhSn*(2*AcpNO*₂*Ph*)*Cl*₂] (6)

The organotin(IV) complexes were obtained by refluxing an ethanol solution of the desired hydrazone with $[Bu^nSnCl_3]$ or $[PhSnCl_3]$ in 1:1 ligand-to-metal molar ratio. The solids were washed with ethanol followed by diethylether and then dried *in vaccuum*. Crystals of complexes **1**, **2**, **3** and **5** were obtained from their ethanolic solutions and were stable in the air. Slow evaporation of **6** in 1:9 dmso:acetone solution gave crystals of [PhSn(2AcpNO₂Ph)Cl₂]·dmso, (**6a**).

n-Butyl(dichloro)(2-acetylpyridinephenylhydrazonato)tin(IV), [BuⁿSn(2AcPh)Cl₂], (1): Color: Yellow. Yield: 71%. Anal. Calc. (C₁₈H₂₁Cl₂N₃OSn): C, 44.88; H, 4.36; N, 8.66. Found: C, 44.87; H, 4.16; N, 8.96%. Selected IR bands (v_{max} /cm⁻¹): v_{as} (CH₃) 2952 m; v_s(CH₃) 2869 m; v_{as}(CH₂) 2925 m; v_s(CH₂) 2853 m; v(C-H)ar 3060 m; v(C=N) 1599 m; σ_{ac} (CH₂) 1460; $\sigma_{as}(CH_3)$ 1421; $\sigma_{s}(CH_3)$ 1380; $\rho(py)$ 649 m; $\nu(M-N_{inine})$ 492; v(M-O) 392; v(M-C) 349; v(M-Cl) 250; v(M-N_) 228. ¹H NMR (CDCl₂, 200 MHz, ppm): δ 8.62 (d, 1H, H6); 8.34 (d, 2H, H10,14); 8.20 (t, 1H, H4); 8.01(d, 1H, H3); 7.74 (t, 1H, H5); 7.49(m, 3H, H11, 13, H12); 2.82 (s, 3H, H15); 2.33(t, 2H, H16); 2.07(m, 2H, H17); 1.61(m, 2H, H18); 1.05(t, 3H, H19). ¹³C NMR (CDCl₂, 200 MHz, ppm): δ 171.38(C8); 145.13(C6); 144.68(C7); 144.16(C2); 142.49(C4); 132.51(C12); 132.42(C9); 129.00(C10,14); 128.15(C11,13); 127.05(C5); 125.06(C3); 35.50(C16); 27.51(C17); 25.60(C18); 13.68(C19); 13.53(C15). ¹¹⁹Sn NMR (CDCl₃, 200 MHz, ppm): δ –414.39. Molar conductivity (1×10⁻³ mol L⁻¹, dmf): 11.56 Ω⁻¹ cm² mol⁻¹.

n-Butyl(dichloro)(2-acetylpyridine-para-chlorophenylhydrazonato)tin(IV), [BuⁿSn(2AcpClPh)Cl₂], (2): Color: Yellow. Yield: 83%. Anal. Calc. (C₁₈H₂₁Cl₃N₃OSn): C, 41.62; H, 3.88; N, 8.09. Found: C, 41.77; H, 3.41; N, 8.39%. Selected IR bands (v_{max} /cm⁻¹): v_{as} (CH₃) 2957 m; $v_{e}(CH_{2})$ 2870 m; $v_{e}(CH_{2})$ 2928 m; $v_{e}(CH_{2})$ 2859 m; v(C-H)ar 3081 m; v(C=N) 1599 m; σ_{as} (CH₂) 1448; σ_{as} (CH₃) 1448; $\sigma_s(CH_3)$ 1386; $\rho(py)$ 647 m; $\nu(M-N_{imine})$ 481; $\nu(M-O)$ 402; v(M-C) 360; v(M-Cl) 250; v(M-N_{nv}) 228. ¹H NMR (CDCl₃, 200 MHz, ppm): δ 8.63(d, 1H, H6); 8.24(d, 2H, H10,14); 8.20(m, 1H, H4); 8.02(d, 1H, H3); 7.67(t, 1H, H5); 7.44(d, 2H, H11,13); 2.81(s, 3H, H15); 2.32(t, 2H, H16); 2.08(m, 2H, H17); 1.60(m, 2H, H18); 1.05(s, 3H, H19). ¹³C NMR (CDCl₂, 200 MHz, ppm): δ 170.50 (C8); 145.28(C6); 145.05(C7); 144.14(C2); 142.56(C4); 138.66(C12); 130.98(C9); 130.43(C11,13); 128.13(C10,14); 127.26(C5); 125.20(C3); 35.60(C16); 27.52(C17); 25.61(C18); 13.68(C19); 13.56(C15). 119Sn NMR (CDCl₂, 200 MHz, ppm): δ –413.66. Molar conductivity (1×10⁻³ mol L⁻¹, dmf): 14.33 Ω^{-1} cm² mol⁻¹.

n-Butyl(dichloro)(2-acetylpyridine-para-nitrophenylhydrazonato)tin(IV), [BuⁿSn(2AcpNO₂Ph)Cl₂], (3): Color: Yellow. Yield: 74%. Anal. Calc. $(C_{18}H_{20}Cl_2N_4O_3Sn)$: C, 40.79; H, 3.80; N, 10.57. Found: C, 40.92; H, 3.72; N, 10.48%. Selected IR bands (v_{max}/cm^{-1}): $v_{as}(CH_3)$ 2956 m; $v_{s}(CH_{2})$ 2869 m; $v_{as}(CH_{2})$ 2927 m; $v_{s}(CH_{2})$ 2858 m; v(C-H)ar 3077 m; v(C=N) 1596 m; σ_{as} (CH₂) 1456; σ_{as} (CH₃) 1443; $\sigma_s(CH_3)$ 1384; $\rho(py)$ 649 m; $\nu(M-N_{imine})$ 479; $\nu(M-O)$ 394; v(M-C) 364; v(M-Cl) 249; v(M-N_w) 214. ¹H NMR (CDCl₂, 200 MHz, ppm): δ 8.68(d, 1H, H6); 8.49(d, 2H, H10,14); 8.30(m, 2H, H11,13); 8.28(m, 1H, H4); 8.11(d, 1H, H3); 7.84(t, 1H, H5); 2.87(s, 3H, H15); 2.34(t, 2H, H16); 2.05(m, 2H, H17); 1.61(m, 2H, H18); 1.05(t, 3H, H19). ¹³C NMR (CDCl₃ 200 MHz, ppm): 169.36(C8); 150.00(C7); 146.89(C2); 145.51(C6); 143.93(C12); 142.62(C4); 138.41(C9); 129.85(C10,14); 127.60(C5); 125.45(C3); 123.21(C11,13); 35.79(C16); 27.52(C17); 25.61(C18); 13.73(C19); 13.68(C15). 119Sn NMR $(CDCl_2, 200 \text{ MHz}, \text{ppm}): \delta -412.33$. Molar conductivity $(1 \times 10^{-3} \text{ mol } \text{L}^{-1}, \text{ dmf})$: 10.66 $\Omega^{-1} \text{ cm}^2 \text{ mol}^{-1}$.

 $\label{eq:Phenyl} Phenyl(dichloro)(2-acetylpyridine-phenylhydrazonato) tin(IV), [PhSn(2AcPh)Cl_2], (4): Color: Yellow. Yield: 85\%. Anal. Calc. (C_{20}H_{17}Cl_2N_3OSn): C, 47.57; H, 3.39; N, 8.32. Found: C, 46.66; H, 3.06; N, 8.80%. Selected$

IR bands (ν_{max} /cm⁻¹): v(C-H)ar 3061 m; v(C=N) 1600 m; ρ(py) 650 m; v(M-N_{imine}) 494; v(M-C) 441; v(M-O) 395; v(M-Cl) 242; v(M-N_{py}) 229. ¹H NMR (CDCl₃, 200 MHz, ppm) main signals: δ 8.41(m, 1H, H4); 8.40(m, 1H, H6); 8.33(d, 2H, H17,21); 8.21(d, 2H, H10,14); 8.04(d, 1H, H3); 7.60(m, 1H, H5); 7.59(m, 3H, H18,19,20); 7.50(m, 1H, H12); 7.44(m, 2H, H11,13); 2.88(s, 3H, H15). ¹³C NMR (CDCl₃, 200 MHz, ppm) main signals: δ 171.53(C8); 145.06(C6); 144.11(C2); 142.61(C4); 139.92(C16); 136.62(C17,21); 132.59(C12); 132.16(C9); 131.13(C19); 129.08(C10,14,18,19); 128.17(C11,13); 126.99(C5); 124.04(C3); 13.45(C15). ¹¹⁹Sn NMR (CDCl₃, 200 MHz, ppm): δ –443.37. Molar conductivity (1×10⁻³ mol L⁻¹, dmf): 20.51 Ω⁻¹ cm² mol⁻¹.

Phenyl(dichloro)(2-acetylpyridine-para-chlorophenylhydrazonato)tin(IV), [PhSn(2AcpClPh)Cl₂] (5): Color: Yellow. Yield: 86%. Anal. Calc. (C₂₀H₁₄Cl₂N₂OSn): C, 44.53; H, 2.99; N, 7.99. Found: C, 43.57; H, 2.82; N, 8.04%. Selected IR bands (v_{max} /cm⁻¹): v(C-H)ar 3050 m; ν (C=N) 1597 m; ρ (py) 647 m; ν (M-N_{imine}) 476; ν (M-C) 443; v(M-O) 403; v(M-Cl) 260; v(M-N_n)233. ¹H NMR $(CDCl_2, 200 \text{ MHz}, \text{ppm}): \delta 8.42(\text{m}, 1\text{H}, \text{H6}); 8.40(\text{m}, 1\text{H}, \text{H6})$ H4); 8.27(d, 2H, H17, 21); 8.21(d, 2H, H11, 13); 8.06(d, 1H, H3); 7.63(t, 1H, H5); 7.59(m, 3H, H18, 19, 20); 7.41(d, 2H, H10,14); 2.88(s, 3H, C15). ¹³C NMR (CDCl₂, 200 MHz, ppm): δ 170.58(C8); 145.11(C6); 144.46(C7); 144.07(C2); 142.77(C4); 139.75(C16); 138.76(C12); 136.50(C17,21); 131.22(C19); 131.00(C9); 130.47(18,20); 129.13(C11,13); 128.50(C10,14); 127.12(C5); 125.04(C3); 13.47(C15). ¹¹⁹Sn NMR (CDCl₂, 200 MHz, ppm): δ –442.65. Molar conductivity (1×10⁻³ mol L⁻¹, dmf): 20.53 Ω^{-1} cm² mol⁻¹.

Phenyl(dichloro)(2-acetylpyridine-para-nitrophenylhydrazonato)tin(IV), [PhSn(2AcpNO_Ph)Cl_], (6): Color: Yellow. Yield: 82%. Anal. Calc. $(C_{20}H_{16}Cl_2N_4O_3Sn)$: C, 43.68; H, 2.93; N, 10.19. Found: C, 43.23; H, 3.00; N, 10.28%. Selected IR bands (v_{max} /cm⁻¹): v(C-H)ar 3058 m; ν (C=N) 1596 m; ρ (py) 649 m; ν (M-N_{imine}) 486; v(M-C) 450; v(M-O) 394; v(M-Cl) 250; v(M-N_{nv}) 231. ¹H NMR (CDCl₂, 200 MHz, ppm): δ 8.50(d, 2H, H11,13); 8.42(m, 1H, H6); 8.41(m, 1H, H4); 8.28(m, 2H, H17,21); 8.25(m, 2H; H10,14); 8.13(d, 1H, H3); 7.70(t, 1H, H5); 7.61(m, 3H, H18,19,20); 2.93(s, 3H; H15). ¹³C NMR (CDCl₂, 200 MHz, ppm): δ 169.41(C8); 150.06(C7); 146.32(C2,C12); 145.28(C6); 142.79(C4); 139.48(C16); 138.43(C9); 136.45(C17,21); 131.36(C19); 129.99(C18,20); 129.21(C11,13); 127.56(C10,14); 125.38(C5); 123.27(C3); 13.64(C15). ¹¹⁹Sn NMR (CDCl₃, 200 MHz, ppm): δ –441.42. Molar conductivity $(1 \times 10^{-3} \text{ mol } \text{L}^{-1}, \text{ dmf})$: 18.75 $\Omega^{-1} \text{ cm}^2 \text{ mol}^{-1}$.

X-ray crystallography

Crystals of [BuⁿSn(2AcPh)Cl₂] (1), [BuⁿSn(2AcpClPh) Cl₂] (2), [BuⁿSn(2AcpNO₂Ph)Cl₂] (3), [PhSn(2AcPh)Cl₂] (4) and [PhSn(2AcpNO₂Ph)Cl₂]•dmso (6a) were mounted on glass fibers and used for data collection. X-ray diffraction data collection was performed on a Oxford-Diffraction *GEMINI* diffractometer (LabCri) using graphite-Enhance Source MoK_{α} radiation ($\lambda = 0.71069$ Å) at 293(2) K. Data integration and scaling of the reflections were performed with the Crysalis suite.³⁴ Final unit cell parameters were based on the fitting of all reflections positions. The structures were solved by direct methods using the program SHELXS-97³⁵ and refined by full-matrix least-squares techniques against F² using SHELXL-97.³⁶ Positional and anisotropic atomic displacement parameters were refined for non hydrogen atoms, except for those involved in disorder systems for which isotropic ADP was used. Although some hydrogen atoms could be identified in a Fourier difference map, in the final model they were geometrically positioned and refined using a riding model. Criteria of a satisfactory complete analysis were the ratios of rms shift to standard deviation less than 0.001 and no significant features in final difference maps. Molecular graphics were obtained from ORTEP.³⁷ A summary of the crystal's data, experimental details and refinement results are listed in Table 1.

In vitro antimicrobial activity

Antimicrobial activity was evaluated by minimum inhibitory concentration (MIC) using the macrodilution test.³⁸ *S. aureus* ATCC6538 stored in Brain Heart Infusion

Table 1. Crystal data and refinement results for $[Bu^nSn(2AcPh)Cl_3]$ (1), $[Bu^nSn(2AcpClPh)Cl_3]$ (2), $[Bu^nSn(2AcpNO_2Ph)Cl_3]$ (3), $[PhSn(2AcPh)Cl_3]$ (4) and $[PhSn(2AcpNO_2Ph)Cl_3]$ -dmso (6a)

Compound		1	2	3	4	6a
Empirical Formula		C ₁₈ H ₂₁ Cl ₂ N ₃ OSn	C ₁₈ H ₂₀ Cl ₃ N ₃ OSn	$C_{18}H_{20}C_{12}N_4O_3Sn$	C ₂₀ H ₁₇ C ₁₂ N ₃ OSn	C ₂₂ H ₂₂ Cl ₂ N ₄ O ₄ SSn
Formula Weight		484.97	519.41	529.97	504.96	628.09
Crystal System		orthorhombic	orthorhombic triclin		triclinic	monoclinic
Space Group		<i>F</i> d d 2	<i>F</i> d d 2	P-1	<i>P</i> -1	$P2_{1}/c$
Unit cell dimensions	<i>a</i> , Å	27.440(4)	26.6226(17)	12.9843(4)	11.4662(3)	14.4967(4)
	b, Å	22.024(4)	23.9627(18)	13.3554(6)	12.5545(3)	13.4617(4)
	<i>c</i> , Å	13.316(2)	13.3438(7)	14.1000(7)	16.2705(4)	13.3607(3)
	α, °	90	90	111.487(4)	112.147(2)	90
	β, °	90	90	94.407(3)	95.650(2)	102.523(3)
	γ, °	90	90	105.649(3)	101.670(2)	90
Volume, Å ³		8047(2)	8512.7(10)	2148.31(16)	2084.75(9)	2545.31(12)
Z / Density calc., Mg/n	n ³	16 / 1.601	16 / 1.621	4 / 1.639	4 / 1.609	4 / 1.639
Absorption coefficient, mm ⁻¹		1.546	1.589	1.464	1.496	1.332
F(000)		3872	4128	1056	1000	1256
Crystal size, mm		$0.27 \times 0.24 \times 0.12$	$0.29 \times 0.24 \times 0.18$	$0.28 \times 0.25 \times 0.10$	$0.24 \times 0.18 \times 0.09$	$0.19 \times 0.19 \times 0.05$
θ range for data collection, °		2.86 to 26.37	3.07 to 26.37	2.76 to 26.37	2.61 to 26.37	2.79 to 26.37
Index range, θ	Index range, θ		$-24 \le h \le 33$	$-16 \leq h \leq 16$	$-14 \leq h \leq 11$	$-18 \leq h \leq 18$
		$-27 \le k \le 23$	$-29 \le k \le 29$	$-13 \le k \le 16$	$-14 \le k \le 15$	$-16 \le k \le 16$
	< 27 a	$-15 \le l \le 16$	$-13 \le l \le 16$	$-17 \le l \le 16$	$-20 \le l \le 19$	$-16 \le l \le 13$
Completeness to $\theta = 26.37^{\circ}$		99.9%	99.6%	99.9%	99.9%	99.9%
Absorption correction		none	none	none	none	none
Goodness-of-fit on F^2		0.927	1.072	0.814	0.869	0.829
Reflections collected/unique (R_{int})		12047 / 3852 (0.0647)	6718 / 3555 (0.0319)	19702 / 8769 (0.0474)	18566 / 8525 (0.0295)	14702 / 5212 (0.0447)
Absolute structure parameter		0.03(5)	0.06(5)	-	-	-
Data / restraints / parameters		3852 / 11 / 224	3555 / 6 / 235	8769 / 8 / 470	8525 / 0 / 487	5212 / 0 / 307
Observed reflections, $I > 2\sigma(I)$		2232	2405	4288	5929	3261
Final <i>R</i> indices $[I > 2\sigma(I)]$		$R_1 = 0.0477,$ $wR_2 = 0.1077$	$R_1 = 0.0483,$ $wR_2 = 0.1335$	$R_1 = 0.0442,$ $wR_2 = 0.0963$	$R_1 = 0.0257,$ $wR_2 = 0.0441$	$R_1 = 0.0324,$ $wR_2 = 0.0543$
R indices (all data)		$R_1 = 0.0879,$ $wR_2 = 0.1166$	$R_1 = 0.0740,$ $wR_2 = 0.1460$	$R_1 = 0.1028,$ $wR_2 = 0.1055$	$R_1 = 0.0469,$ $wR_2 = 0.0463$	$R_1 = 0.0655,$ $wR_2 = 0.0580$
Larg. peak and hole, e $Å^{-3}$		1.047 / -0.660	0.739 /-1.179	0.951 / -0.490	0.309 / -0.279	0.968 / -0.569

(BHI) broth were subcultured for testing in the same medium and grown at 37 °C. Then the bacterial cells were suspended in BHI, according to Clinical and Laboratory Standards Institute (CLSI) guidelines,³⁹ to produce a suspension of about 10⁵ CFU (colony forming units) mL⁻¹. *C. albicans* ATCC18804 stored in Sabouraud broth was sub-cultured for testing in the same medium and grown at 37 °C. Then the yeast cells were suspended according to the McFarland protocol³⁹ in saline solution to produce a suspension of about 10³ cells mL⁻¹.

For both the antibacterial and antifungal activity evaluations serial dilutions of the compounds, previously dissolved in dmso, were prepared in test tubes. A 24-h old inoculum (100 μ L) was added to each tube. The MIC, defined as the lowest concentration of the test compound which inhibits the visible growth after 20 h, was determined after incubation at 37 °C. Tests using tetracycline (in the case of *S. aureus*) or fluconazole (in the case of *C. albicans*) as reference and dmso as negative control were carried out in parallel. The final dmso concentration never exceeded 1%. In all cases no inhibition was observed with 5% v/v dmso. All tests were performed in triplicates with full agreement between the results.

Results and Discussion

Microanalyses and molar conductivity data suggest the formation of the following complexes: $[Bu^nSn(2AcPh)Cl_2]$ (1), $[Bu^nSn(2AcpClPh)Cl_2]$ (2), $[Bu^nSn(2AcpNO_2Ph)Cl_2]$ (3), $[PhSn(2AcPh)Cl_2]$ (4), $[PhSn(2AcpClPh)Cl_2]$ (5) and $[PhSn(2AcpNO_2Ph)Cl_2]$ (6), in which an anionic hydrazone is attached to the metal centre along with a *n*-butyl or a phenyl group and the remaining coordination sites are occupied by two chloride ions. X-ray diffraction analysis confirms the structures of 1-4 and 6 (as its dmso solvate, $[PhSn(2AcpNO_2Ph)Cl_2]$ ·dmso, 6a).

Spectroscopic characterization

Infrared spectra

The v(N-H) and v(C=O) absorptions at 3180-3287 cm⁻¹ and 1655-1676 cm⁻¹ in the spectra of the uncomplexed hydrazones were not found in the spectra of complexes (1-6), in accordance with the presence of anionic ligands in the products.

The v(C=C) + v(C=N) composed mode observed at 1590-1460 cm⁻¹ in the spectra of the hydrazones undergoes small shifts in the spectra of the complexes. However a new v(C=N) absorption due to the formation of a second C=N bond was observed at 1625-1632 cm⁻¹ in the spectra of **1-6**, indicating coordination of an anionic hydrazone.

The pyridine in-plane deformation mode at 620-625 cm⁻¹ in the spectra of the hydrazones shifts to 647-650 cm⁻¹ in those of the complexes, suggesting coordination of the heteroaromatic nitrogen.²⁹⁻³¹ In addition, new absorptions at 495-475 cm⁻¹ and 450-350 cm⁻¹ have been attributed to v(Sn-N_{imine}) and v(Sn-C) respectively; absorptions at 230-215 cm⁻¹ were assigned to v(Sn-N_{py}),^{25,28,40} and bands in the 405-390 cm⁻¹ range have been assigned to v(Sn-O). An absorption at 260-250 cm⁻¹ has been attributed to v(Sn-Cl).^{25,28,40} Therefore, the infrared data for the complexes indicate coordination of the hydrazones through the N_{py}-N-O chelating system, as confirmed by the X-ray structure determinations for **1-4** and **6a**.

NMR spectra

The ¹H and ¹³C NMR spectra of the hydrazones were recorded in dmso- d_6 as well as in CDCl₃ except in the case of H2AcpNO₂Ph, which is insoluble in CDCl₃. The spectra of complexes **1-6** were recorded only in CDCl₃ since some of them interact with dmso- d_6 .

The ¹H resonances were assigned on the basis of chemical shifts, multiplicities and coupling constants. The carbon type (C, CH) was determined by using distortionless enhancement by polarization transfer (DEPT135) experiments. The assignments of the protonated carbons were made by 2D heteronuclear multiple quantum coherence experiments (HMQC). A ¹¹⁹Sn NMR study was performed for all complexes.

The NMR spectra of H2AcPh in CDCl₃ show only one signal for each hydrogen and each carbon, indicating the presence of only one configurational isomer. The signal of N3-H was observed at δ 9.22, suggesting the presence of the *E* isomer. The spectra recorded after 24 h do not show any changes. The spectra recorded in dmso-*d*₆ show two signals for each hydrogen and carbon, in accordance with the presence of the *E* and *Z* isomers (see Figure 1) in solution.²⁹⁻³¹ The signals of N3-H at δ 10.90 and δ 15.80 were attributed to the *E* and *Z* isomers respectively (97% of *E* and 3% of *Z*). The high frequency signal of the *Z* isomer is characteristic of N3-H hydrogen-bonded to the heteroaromatic nitrogen while the signal at δ 10.90 is characteristic of N3-H hydrogen-bonded to the solvent, as previously observed by us.²⁹⁻³¹

H2AcpNO₂Ph is soluble in dmso- d_6 but insoluble in CDCl₃. In the ¹H and ¹³C NMR spectra (dmso- d_6) of H2AcpNO₂Ph only one signal has been observed for each hydrogen and each carbon. The signal of N3-H was found at δ 11.22, suggesting the presence of the *E* configurational isomer, in which N3-H is hydrogen-bonded to the solvent.²⁹⁻³¹ The ¹H and ¹³C NMR spectra of H2AcpClPh in dmso- d_6 show duplicated signals for the hydrogens, in accordance with the presence of two configurational isomers. The signals of N3-H at δ 10.96 and δ 15.86 were attributed to the *E* (98%) and *Z* (2%) isomers, respectively.²⁹⁻³¹ Due to the low solubility of H2AcpClPh in dmso- d_6 only some of the signals of hydrogens could be attributed and no attribution was possible for the carbons of the *Z* isomer.

The spectra of this compound in CDCl_3 also show duplicated signals. However in this case the spectra are characteristic of the presence of the keto and enol forms.⁴¹ In fact no signal of N3-H was observed for the first isomer, according to the presence of the enol form. In addition, the signal of C-O was observed at δ 167.44. The resemblance in position of the carbon signals of H2AcpClPh with those of H2AcPh in the same solvent suggests that the enol isomer adopts the *E* configuration. As an example, the signal of C15 (acetyl) was observed at δ 11.36, close to the signal for the same hydrogen at δ 10.89 observed for the *E* isomer of H2AcPh in CDCl₃. The keto isomer exhibits the signal of N3-H at δ 15.83, characteristic of the *Z* configuration, and the signal of C=O at δ 163.40.

Only one signal was observed for each hydrogen and each carbon in the spectra of complexes **1-6**, suggesting the presence of only one configurational isomer. The signals of C=N, C=O and the pyridine carbons undergo significant shifts in the complexes indicating coordination through the N_{py} –N–O chelating system. Hence the hydrazones adopt the *E* configuration in the complexes, as confirmed by crystal structure determinations of complexes **1-4** and **6a**. Moreover, the signal of N3-H was not observed in the ¹H NMR spectra of the complexes, according to deprotonation of the hydrazones upon coordination.

In the spectra of complexes **1-3**, new signals at δ 2.35-1.05 and at δ 35.80-13.60 were attributed to the hydrogens and carbons, respectively, of the *n*-butyl group attached to the metal.²⁸ In the spectra of complexes **4-6**, the signals at δ 8.33-7.59 and at δ 136.62-129.08 were attributed to the hydrogens and carbons, respectively, of the phenyl group coordinated to tin.⁴⁰

In the ¹¹⁹Sn NMR spectra of complexes **1-6** (CDCl₃) only one signal was observed at $\delta - 414$ (**1**), $\delta - 413$ (**2**), $\delta - 412$ (**3**), $\delta - 443$ (**4**), $\delta - 442$ (**5**) and $\delta - 441$ (**6**) ppm, compatible with the presence of one tin site in solution. The values of the ¹¹⁹Sn signals are at the boundary between six- and seven-coordinated tin(IV) as reported in the literature.⁴² However, considering that the spectra were recorded in a non-coordinating solvent (CDCl₃), only six-coordinated species must be present in solution. The shielding effect on ¹¹⁹Sn of the chloride ions could explain the lower frequencies observed for the ¹¹⁹Sn signals with respect to that expected for a six-coordinated monorganotin(IV).⁴²

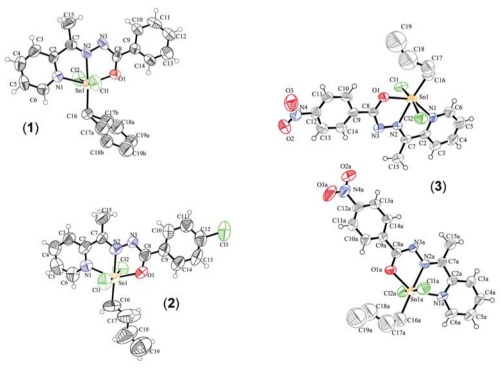


Figure 2. Molecular plot of the asymmetric units of $[(Bu^n)Sn(2AcPh)Cl_2]$ (1), $[(Bu^n)Sn(2AcpClPh)Cl_2]$ (2) and $[(Bu^n)Sn(2AcpNO_2Ph)Cl_2]$ (3) showing the labeling scheme of the non-H atoms and their displacement ellipsoids at the 50% probability level.



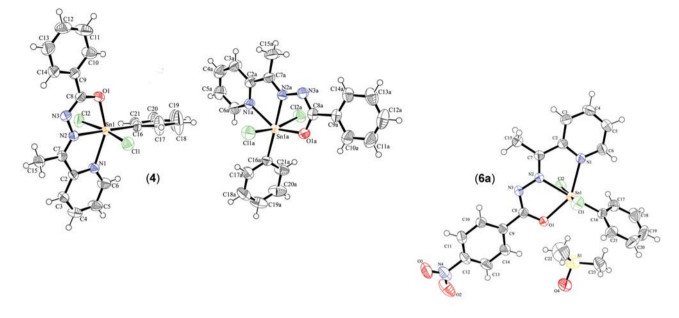


Figure 3. Molecular plot of the asymmetric units of $[PhSn(2AcPh)Cl_2]$ (4) and $[PhSn(2AcpNO_2Ph)Cl_2]$ -dmso (6a) showing the labeling scheme of the non-H atoms and their displacement ellipsoids at the 50% probability level.

X-ray diffraction analysis

Figures 2 and 3 are perspective views of [BuⁿSn(2AcPh)Cl₂] (1), [BuⁿSn(2AcpClPh)Cl₂] (2), [BuⁿSn(2AcpNO₂Ph)Cl₂] (3), [PhSn(2AcPh)Cl₂] (4) and [PhSn(2AcpNO₂Ph)Cl₂]-dmso (**6a**). Selected intramolecular bond distances and angles of the complexes are given in Tables 2 and 3. The distances and angles in all complexes are similar. In all complexes tin(IV) is coordinated to a tridentate anionic ligand, a *n*-butyl (1-3) or a phenyl (4 and **6a**) group and two chloride ions *trans* to each other in an distorted octahedral environment. In the crystal structures of **3** and **4**, two symmetrically independent molecules were found in the asymmetric unit (see Figure 2 and Figure 3).

In complexes **1-4** and **6a** the N1–Sn–O1 angle of *ca*. 144.3-146.7° deviates markedly from the ideal value of 180°, probably due to the spatial requirements of the ligand chelating system. By contrast, the C11–Sn–C12 and N2–Sn–C16 angles, which do not involve sterical hindrance, are approximately 167.3-171.4° and 172.5-179.1° (see Table 3). In all complexes the metal atom lies in the plane calculated for the N1N2O1C16 atoms. In structures **1-3** the distance of the tin atom to the calculated plane varies from 0.023Å to 0.059 Å; in structures **4** and **6a** this distance is smaller (0.002 to 0.009 Å).

In all structures the hydrazone skeleton (C7(C15) N2N3C8O1) is planar, with medium deviation of the fitted atoms from the calculated plane ranging from 0.0055 to 0.0266 Å. In **1-4** and **6a** the hydrazone adopts the *E* configuration in relation to the C7=N2 double bond. The

angle between the pyridine and the hydrazone planes varies from 1.5(6) to $7.5(3)^{\circ}$, while the angle between the hydrazone and the phenyl planes varies in the 0.5(2) to $14.9(2)^{\circ}$ range. Therefore the hydrazone ligand in all complexes is almost planar. Complexes **3** and **6a** show the highest deviation from planarity for the angle between the hydrazone chain and the phenyl ring.

The molecular structure of H2AcPh has been determined previously by other authors.⁴³ The hydrazone adopts the EZ conformation in relation to the C7=N2 and N3-C8 bonds. The expected lengthening of the C8-O1 bonds from 1.219(2) Å in H2AcPh⁴³ to 1.279(11) Å in 1 and 1.292(3) and 1.302(3) Å in 4, together with the shortening of the N3-C8 bond from 1.347(3) Å in H2AcPh⁴³ to 1.337(12) Å in **1** and 1.321(3) and 1.327(3) Å in 4, were observed (see Table 2). Therefore the C8-O1 bond changes from a double to a predominantly single bond whereas N3-C8 acquires some double bond character. The bond distances between tin and the imine nitrogen in the complexes, 2.184(4)-2.161(7) Å, are slightly shorter than the distance between tin and the pyridine nitrogen, 2.288(4)-2.2354(18) Å. The Sn-O bond lengths lie in the 2.0916(17)-2.108(3) Å range.

Bond distances in the five complexes are similar to those found in other tin(IV) complexes with hydrazones, as in *trans*-dimethyl[2,6-diacetylpyridinebis(2-theoylhydrazone)] tin(IV)⁴⁴, where N3–C8 and C8–O1 bond distances of 1.322(8) Å and 1.286(7) Å, respectively, and Sn–C, Sn–N_{py}, Sn–N_{imine} and Sn–O bond distances of 2.102(6), 2.327(4), 2.255(5) and 2.209(4) Å, respectively, were observed.

A twisting of approximately 180° in the C2–C7 bond of the hydrazone to match the steric requirements of tridentate coordination was evidenced. Hence some angles undergo significant changes on complexation. The C8–N3–N2 angle goes from 117.02(13)° in H2AcPh⁴³ to 110.7(7)° in **1** and to 109.5(2)° and 109.8(2)° in **4**; N3–C8–O goes from 123.32(15)° in the free base⁴³ to 125.7(8)° in **1** and to 126.4(2)° and 125.2(2)° in **4**.

The molecular packings of all complexes show weak interactions involving Cl, H and O atoms with formation of molecular chains as shown in Figure 4. **1-4** show interactions between adjacent molecules involving $\text{Cl}\cdots\text{H}_{py}$. In **4** and **6a** short Cl \cdots H contacts are observed involving chlorides and hydrogens from the hydrazone phenyl ring. In **6a** Cl \cdots H interactions were observed as well, along with short H \cdots O contacts involving the oxygens from dmso and from the nitro group and hydrogens from pyridine and dmso.

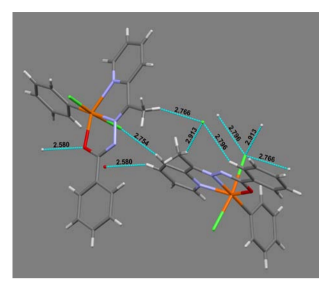


Figure 4. Molecular structure scheme of $[PhSn(2AcPh)Cl_2]$ (4) showing short intermolecular contacts between $Cl \cdots H$ and $O \cdots H$ atoms.

Table 2. Selected bond lengths (Å) for $[Bu^nSn(2AcPh)Cl_2]$ (1), $[Bu^nSn(2AcpClPh)Cl_2]$ (2), $[Bu^nSn(2AcpNO_2Ph)Cl_2]$ (3), $[PhSn(2AcPh)Cl_2]$ (4) and $[PhSn(2AcpNO_2Ph)Cl_2]$ -dmso (6a)

Bond	1	2	:	3		4	6a
C2C7	1.495(11)	1.504(13)	1.456(7)	1.483(7)	1.487(3)	1.467(4)	1.478(4)
C7-C15	1.505(12)	1.474(12)	1.484(7)	1.474(7)	1.479(3)	1.486(3)	1.479(4)
C7-N2	1.253(11)	1.298(10)	1.283(6)	1.279(6)	1.285(3)	1.285(3)	1.282(4)
N2-N3	1.381(9)	1.363(10)	1.381(6)	1.390(5)	1.381(2)	1.369(3)	1.375(3)
N3-C8	1.337(12)	1.305(11)	1.346(6)	1.308(7)	1.321(3)	1.327(3)	1.313(4)
C8–C9	1.455(12)	1.475(13)	1.471(7)	1.480(7)	1.490(3)	1.470(3)	1.487(4)
C8-O1	1.279(11)	1.309(11)	1.275(6)	1.305(6)	1.292(3)	1.302(3)	1.293(3)
Sn1–O1	2.108(6)	2.105(7)	2.096(4)	2.108(3)	2.0996(14)	2.0916(17)	2.104(2)
Sn1–N1	2.246(8)	2.280(8)	2.278(4)	2.288(4)	2.2354(18)	2.2484(19)	2.264(3)
Sn1-N2	2.171(7)	2.161(7)	2.184(4)	2.175(4)	2.165(2)	2.1713(18)	2.170(3)
Sn1-Cl1	2.456(2)	2.460(2)	2.445(2)	2.445(2)	2.4584(7)	2.4705(7)	2.4771(9)
Sn1–Cl2	2.494(3)	2.455(3)	2.500(2)	2.473(2)	2.4532(7)	2.4471(7)	2.4452(9)
Sn1-C16	2.119(9)	2.166(12)	2.140(7)	2.137(8)	2.138(3)	2.124(2)	2.135(3)

Table 3. Selected angles (°) for $[Bu^nSn(2AcPh)Cl_2]$ (1), $[Bu^nSn(2AcpClPh)Cl_2]$ (2), $[Bu^nSn(2AcpNO_2Ph)Cl_2]$ (3), $[PhSn(2AcPh)Cl_2]$ (4) and $[PhSn(2AcpNO_2Ph)Cl_3]$ -dmso (6a)

Angle	1	2		3		4	6a
C2-C7-N2	116.4(7)	114.1(8)	114.5(5)	113.6(5)	114.0(2)	115.0(2)	114.7(3)
N2-C7-C15	125.7(8)	123.9(8)	123.4(5)	124.6(5)	124.0(2)	122.5(2)	123.6(3)
C7-N2-N3	123.0(7)	120.2(7)	121.5(4)	119.7(4)	121.5(2)	122.2(2)	121.7(3)
C8-N3-N2	110.7(7)	111.2(7)	109.6(4)	109.3(4)	109.5(2)	109.8(2)	110.1(3)
N3-C8-C9	116.6(9)	117.2(9)	115.5(5)	115.5(5)	116.0(2)	117.6(2)	115.9(3)
N3-C8-O1	125.7(8)	124.0(8)	125.5(5)	127.1(5)	126.4(2)	125.2(2)	126.5(3)
C9–C8–O1	117.6(8)	118.8(8)	119.0(4)	117.3(5)	117.7(2)	117.6(2)	117.6(3)
N2-Sn1-C16	175.5(3)	176.1(5)	174.7(2)	172.5(2)	179.06(8)	178.82(9)	176.71(11)
N1-Sn1-O1	146.7(2)	145.3(3)	144.96(16)	144.32(15)	146.12(7)	145.88(6)	145.97(9)
Cl1-Sn1-Cl2	167.76(12)	169.41(13)	168.97(5)	171.36(6)	169.54(2)	167.30(2)	170.75(3)

Antimicrobial activity

The minimum inhibitory concentrations (MICs) of the hydrazones and their organotin complexes against *S. aureus* and *C. albicans* are reported in Table 4. Among the hydrazones the antibacterial activity follows the order H2AcPh < H2AcpNO₂Ph < H2AcpClPh, suggesting that the electron-withdrawing effect of the nitro and chloro groups probably favors the antibacterial effect. The calculated values of log P⁴⁵ were: 1.98 for H2AcPh, 2.58 for H2AcpClPh and 1.92 for H2AcpNO₂Ph. Therefore the higher lipophilicity of H2AcpClPh probably accounts in part for its higher activity.

Table 4. Minimum inhibitory concentrations (MIC) of the hydrazones and their organotin complexes against *Staphylococcus aureus* and *Candida albicans*

Compound	MIC / (µmol mL ⁻¹) C. albicans	MIC / (µmol mL ⁻¹) S. aureus
H2AcPh	0.213	0.428
$[Bu^{n}Sn(2AcPh)Cl_{2}](1)$	0.210	0.255
$[PhSn(2AcPh)Cl_2]$ (4)	0.206	0.132
H2AcpClPh	0.046	0.197
$[Bu^{n}Sn(2AcpClPh)Cl_{2}] (2)$	0.045	0.059
$[PhSn(2AcpClPh)Cl_2] (5)$	0.046	0.057
H2AcpNO ₂ Ph	> 0.179	0.353
[BunSn(2AcpNO2Ph)Cl2] (3)	0.096	0.125
$[PhSn(2AcpNO_{2}Ph)Cl_{2}] (6)$	0.093	0.114
[Bu ⁿ SnCl ₃]	> 0.86	> 1.88
[PhSnCl ₃]	0.158	0.249
Tetracycline	-	0.020
Fluconazole	0.033	-

[BuⁿSnCl₃] proved to be inactive against *S. aureus* while [PhSnCl₃] exhibited antibacterial activity in the assayed concentrations. The hydrazones showed antibacterial activity with MIC values similar to the phenyltin salt. Upon coordination the antibacterial activities of both the hydrazones and tin substantially increase. A synergistic effect involving tin and the hydrazone might be responsible for this enhancement.

The complex of H2AcPh with phenyltin is more active than its *n*-butyltin analogue, but the *n*-butyltin and phenyltin complexes with the other two hydrazones are equally active. The activities of the *n*-butyltin complexes with the different hydrazones follows the order [BuⁿSn(2AcPh)Cl₂] < [BuⁿSn(2AcpNO₂Ph)Cl₂] < [BuⁿSn(2AcpClPh)Cl₂], which is the same order of activity of the free hydrazones. However, the activities of the phenyltin complexes are [PhSn(2AcPh)Cl₂] = $[PhSn(2AcpNO_2Ph)Cl_2] < [PhSn(2AcpClPh)Cl_2].$ The most active complexes were those with H2AcpClPh (2 and 5), with MIC values similar to that determined for tetracycline.

Among the hydrazones H2AcpClPh exhibited the highest activity against *C. albicans*. Upon coordination the antifungal activity of the hydrazones increased in complexes 3 and 6, but remained the same in the other cases. The antifungal activity increased in complexes 2, 3, 5 and 6 relative to the organotin salts.

Like in the case of the antibacterial activity, the best results were obtained for the organotin complexes with H2AcpClPh. The MIC values for complexes 2 and 5 (0.045 μ mol mL⁻¹) were the same, suggesting that the *n*-butyl or the phenyl group in the metal coordination sphere have the same effect. This value of MIC is of the same order of magnitude of that obtained for fluconazole. Likewise, the MIC values are the same for complexes 3 and 6.

Conclusions

Organotin compounds are known for their innumerous bioactivities but are also often rather toxic.^{46,47} Coordination of organotin salts with organic ligands could be an interesting strategy for activity improvement⁴⁸ or toxicity reduction. In the present work we demonstrated that coordination of organotin salts to hydrazones might improve the pharmacological profile of both metal and ligands.⁴⁹

Supplementary Information

Crystallographic data have been deposited with the Cambridge Crystallographic Data Centre (deposition numbers CCDC 747621 (1), CCDC 747620 (2), CCDC 747624 (3), CCDC 747623 (4) and CCDC 747622 (6a)). Copies of available material can be obtained by request to CCDC, 12 Union Road, Cambridge CB2 1EZ, UK (fax 44-1223-336033 or e-mail: deposit@ccdc.cam.ac.uk).

Acknowledgments

This work was supported by FAPEMIG, CNPq and Instituto do Milênio em Inovação e Desenvolvimento de Novos Fármacos e Medicamentos (IM-INOFAR) (Proc. CNPq 573.564/2008-6), Brazil.

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Received: September 14, 2009 Web Release Date: April 12, 2010