

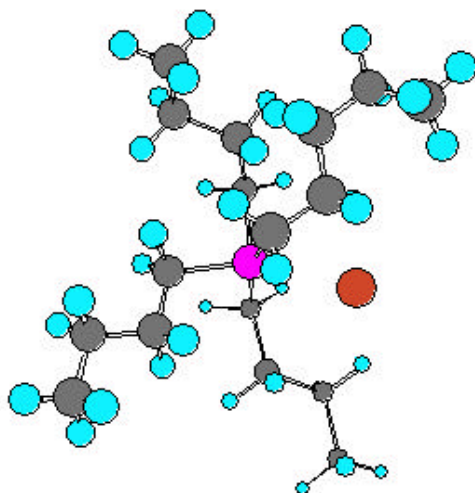
CYTEC

Technology ahead of its time™

www.cytec.com

CYPHOS® IL 163

Phosponium Ionic Liquid



Introduction:

Why consider a “phosphonium” ionic liquid?

Soon after the discovery that certain nitrogen based room temperature liquid salts were found to be useful as battery electrolytes (1a,1b,), interest in these and similar salts as novel fluids and solvents developed. There were a scant number of papers during the 1980s and early 1990s but mainly due to the efforts of the group at The Queen’s University – Belfast, headed up by Professor Ken Seddon, there has been an exponential rise in interest and number of publications in the last 7 to 8 years. (2) Indeed, almost an entire issue of Green Chemistry (3) has been devoted to ionic liquids.

Perhaps one of the most influential publications to direct industrial attention to ionic liquids was a feature article entitled “Designer Solvents” in C&E News – March 30,1998 in which Ken Seddon, Robin Rogers, Tom Welton, Helene Olivier and others elaborated on the potential of ionic liquids. While the article dealt almost entirely with nitrogen based ionic liquids, there was a brief reference by Ken Seddon which alluded to the fact that phosphonium salts are also a potential source of numerous ionic liquids. This brief reference to phosphonium ionic liquids is very much representative of the current fraction of publications relating to phosphonium based ionic liquids. With the exception of several papers and patents by George Parshall in the mid 1970s using stannate and germanate salts and John Knifton et al in the early 1990s which centre on the use of molten tetrabutylphosphonium bromide as an ionic solvent, almost the entire volume of ionic liquid literature deals with nitrogen based systems and in particular, those based on 2-methylimidazolium salts.

There was a good reason for the lack of phosphonium based ionic liquid publications – availability of the starting material! While Cytec has been commercially producing phosphine derivatives since 1971, it was not until 1990 that tributylphosphine was produced on a large commercial scale. Since that time, not only has tetrabutylphosphonium chloride and bromide become available in multi ton scales, many other trialkylphosphines and the corresponding quaternary phosphonium salts are or can be manufactured on a large scale.

The phosphonium cation contains four substituents and the various combinations along with the multitude of various available anions represents an enormous number of possible salts. Even when one restricts the cation to the generic formula – $[PR_3R']^+$, the number is still very large. Of course, not all such phosphonium salts are liquid at room temperature, but by a judicious selection of R and R’ as well as the appropriate anion, there are many phosphonium salts which are in fact liquid at room temperature and many more which fall within the broad general definition of ionic liquids as salts which are low melting – that is less than 100 °C.

There are several reasons why one might consider a phosphonium ionic liquid. The most important one for those contemplating an industrial process is availability and cost. Phosphonium salts can meet both of these demands – already Cytec is manufacturing phosphonium salts on a multi ton scale and because of the high volumes, costs will be relatively low. For commercial products, chemical inventory registration is also part of the availability equation. While, most of the possible phosphonium ionic

liquids are still not registered, several are already listed on EINECS, TSCA, EEC, AICS, PICCS and DSL.

Ionic liquids, in general, are not going to be outrageously expensive, but they will not be in the same league as toluene, 2-hydroxypropane (IPA) or tetrahydrofuran (THF). This means that to be economically viable, they must be chemically as well as thermally very stable for multiple recycle use. Even 0.5 to 1% decomposition can lead to major losses after 10 to 20 cycles. Not only will there be solvent losses but there will also be contamination of the ionic liquid solvent and/or products with decomposition byproducts.

In this regard, phosphonium salts are much more thermally stable than the corresponding ammonium salts and even have an edge on imidazolium salts. This is very important for processes which operate at temperatures greater than 100 °C. In addition to being slightly less thermally stable, the imidazolium cation contains protons which are not entirely inert. They are some what acidic which can result in carbene formation. Phosphonium salts, on the other hand have no such acidic protons.

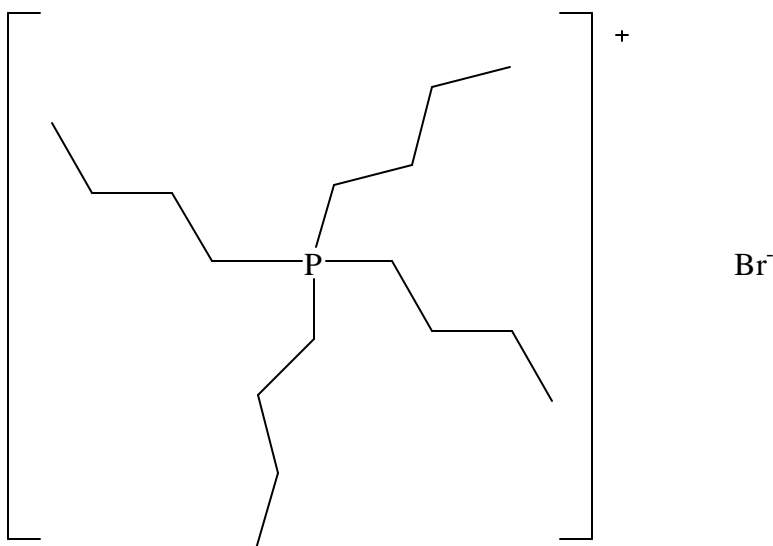
The fact that alkylphosphonium salts are, in general, less dense than water can be beneficial in product work-up steps while decanting aqueous streams which contain inorganic salt byproducts. Imidazolium salts, on the other hand are more dense than water.

Trade Name: **CYPHOS IL 163 phosphonium ionic liquid**

Chemical Name: tetrabutylphosphonium bromide

C.A.S. Number: [3115-68-2]

Registration: EINECS (218-97-0); TSCA; DSL; EEC; PICCS



[CYPHOS IL 163 phosphonium ionic liquid](#) is a crystalline solid melting at 102 °C and might not normally be considered as an “ionic liquid” however, it does come under the broader definition since it is a “low melting” salt. More importantly, it was one of the first to be exhaustively studied for several commercially important industrial processes.

Viscosity:

The viscosity of [CYPHOS IL 163](#) as a function of temperature is presented in Figure 1. Because of the very low viscosity and relatively low molecular weight, [CYPHOS IL 163](#) is well suited as a medium for electrochemistry. These properties are very important considerations when high diffusion coefficients and conductivities are required.

As well as temperature effects, the addition of any substrate will have a dramatic effect on viscosity reduction. This demonstrated also in Figure 1 in which the viscosity/temperature relationship is plotted for 10% solutions of toluene and water.

Thermal Stability:

Typically standard TGA data is reported to indicate the thermal stability of ionic liquids. As long as the experimental conditions are the same – heating rate, atmosphere over the sample and sample size - such data is only a good measure of relative thermal stability.

Using a 10 °C/min heating rate, the onset of weight loss for [CYPHOS IL 163](#) under an air atmosphere is approximately 270 °C while under an atmosphere of dinitrogen, the “apparent” decomposition temperature increases to approximately 310 °C.

However, a more realistic measure of thermal stability is determined by isothermal TGA studies. Under these conditions, the sample is held at various constant temperatures for an extended period of time, during which any weight loss is observed. Figures 2 and 3 depict weight changes for [CYPHOS IL 163](#) under dinitrogen and air atmospheres respectively.

Based on the data in Figures 2 and 3, it would appear that the true upper limit for thermal stability will be approximately 200 °C under a dinitrogen atmosphere. This decreases slightly to about 180 °C under an air atmosphere. This is consistent with most other phosphonium based ionic liquids. It is expected that for most applications, these upper temperature limits will be more than adequate.

Applications:

Prior to there actually being a commercial source of tetra n-butylphosphonium bromide, John Knifton and Jen Jiang Lin published and patented an enormous number of petrochemical applications using this molten salt as a solvent. The applications include the manufacture of ethylene, acetaldehyde, acetic acid, C1 to C4 alcohols, formamides

and ethylene glycol from synthesis gas, methyl acetate from methanol, synthesis of secondary and tertiary amines and homoligation of methanol to ethanol. (4-38)

The above practical industrial applications clearly demonstrate that ionic liquids do not have to be liquid at room temperature. It is only necessary that the salt or more importantly the reaction mixture be liquid at or below the reaction temperature.

More recently, [CYPHOS IL 163](#) has been successfully used as a medium to fluorinate arylchlorides via halide exchange. (39)

Khodaei et. al. have reported high yields of 2-anilinoalkanols from epoxides and analines in a molten tetrabutylphosphonium salt medium (40). In a more recent paper, they also report a high yielding regioselective and enviromentally friendly thiolysis of 1,2-epoxides (41) in a tetrabutylphosphinium salt medium.

Brunet et. al. have reported for the first time the palddium-catalyzed hydroamination of ethylene with aniline in a biphasic system consisting of molten tetrabutylphosphonium bromide and decane (42).

Availability:

[CYPHOS IL 163](#) is now being produced on a multi ton scale and additionally is listed on all the major chemical inventories. Consequently unlimited quantities are now available for developmental and commercial use.

As an alternate, [CYPHOS IL 164](#) - the corresponding chloride salt - is also listed on all the major chemical inventories and is commercially available on a multi ton scale. [CYPHOS IL 164](#) has the added advantage in that it is lower melting (82 °C).

Analysis:

[CYPHOS IL 163](#) typically assays >96% $[\text{PR}_4]^+ \text{Br}^-$ with the main impurities being 0.1 to 0.6% HBr and 0.1 to 1% $[\text{PR}_3\text{H}]^+ \text{Br}^-$. Titration with standardized AgNO_3 in a 75% aqueous 2-hydroxypropane (IPA) solvent will yield the total chloride content. This result when combined with a corresponding NaOH titration will yield the $[\text{PR}_3\text{H}]^+ \text{Br}^-$, HBr and $[\text{PR}_3\text{H}]^+ \text{Br}^-$ content.

^{31}P NMR is generally not suitable for precise analysis. However, the distinctive signal at +33 ppm can be useful for qualitative analysis.

While electro-spray mass spectral analysis (ESMS) is generally not available in every laboratory, this also a very useful tool for both quantitative and qualitative analysis.

Figure 1
CYPHOS IL 163 Viscosity as a Function of Temperature

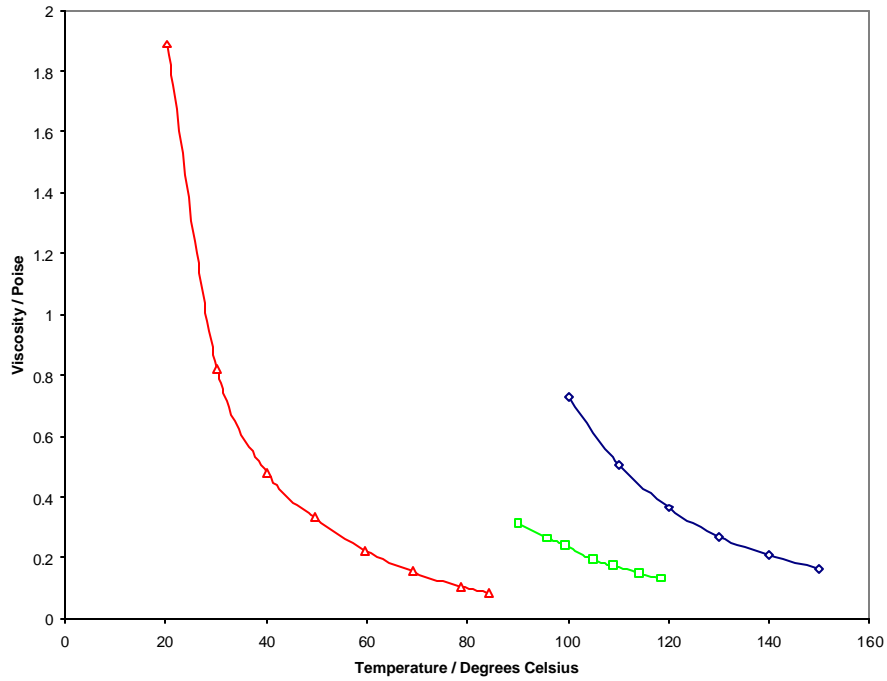


Figure 2:
Isothermal TGA of CYPHOS IL 163 Under a Dinitrogen Atmosphere

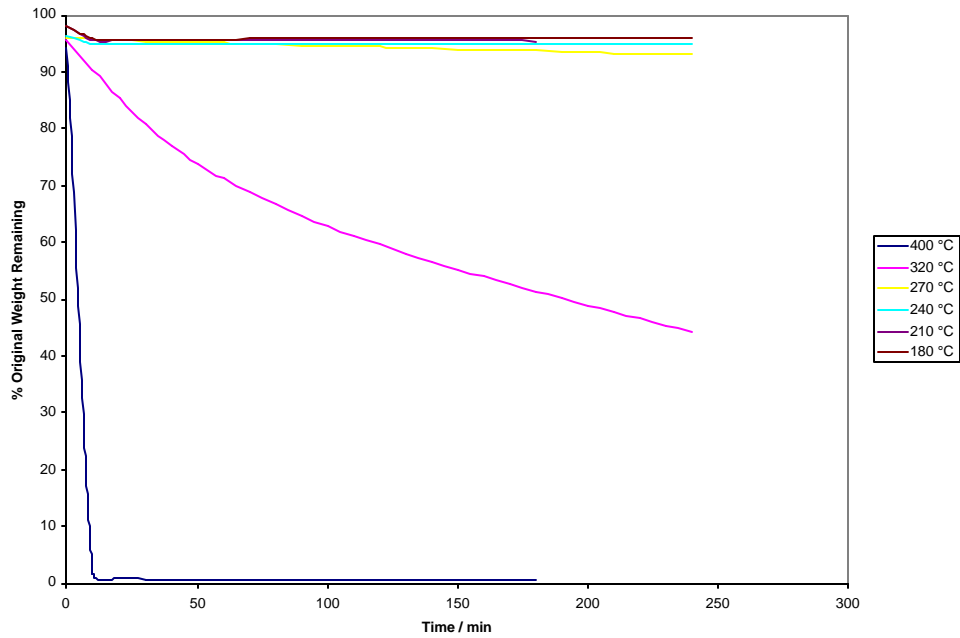
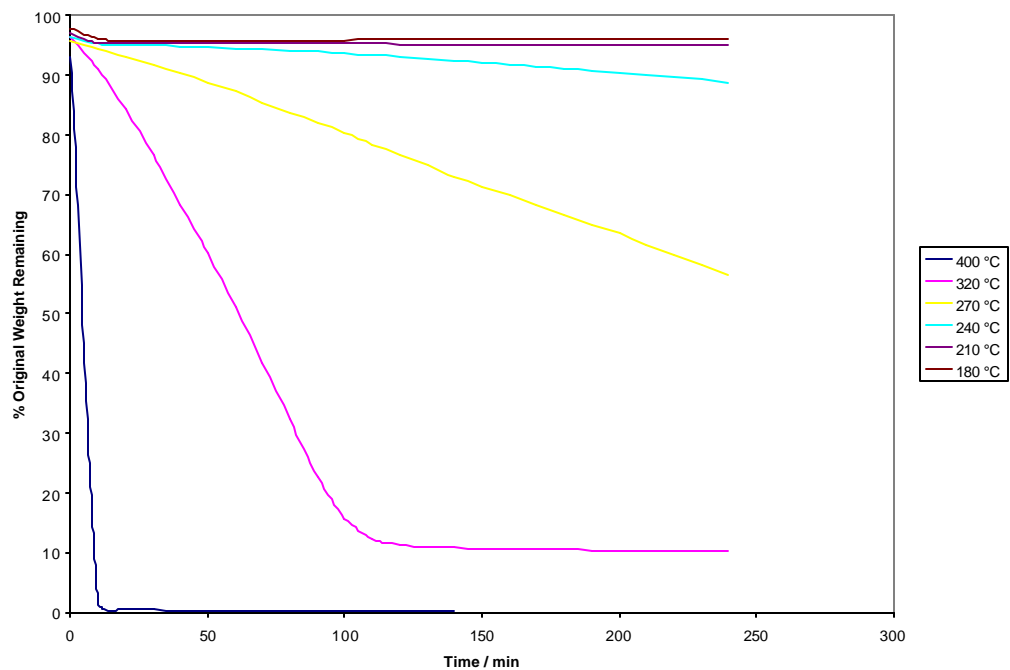


Figure 3:

Isothermal TGA of CYPHOS IL 163 Under an Air Atmosphere



References:

- 1a) Fannin, A. A. Jr; King, L. A.; Levisky, J. A.; Wilkes, J. S.; J. Phys. Chem., 1984, **88**, 2610-2614
- 1b) Fannin, A. A. Jr; Floreani, D. A.; King, L. A.; Landers, J. S.; Piersma, B. J.; Stech, D. J.; Vaughn, R. L.; Wilkes, J. S.; and Williams, J. L., J. Phys. Chem. 1984, **88**, 2614-2621
- 2) "Forward", Professor Ken Seddon, Green Chemistry April, 2002.
- 3) Green Chemistry April, 2002
- 4) Syngas reactions. XII. The selective preparation of acetaldehyde, alcohols, esters and acetic acid from synthesis gas.; Knifton, John F.; Lin, J. J.; Appl. Organomet. Chem. (1989), 3(6), 557-62
- 5) Syngas reactions. Part XIII. The ruthenium 'melt'-catalyzed oxonation of terminal olefins.; Knifton, John F.; Lin, J. J.; J. Mol. Catal. (1988), 47(1), 99-116
- 6) Syn gas reactions. Part XI. The ruthenium melt-catalyzed oxonation of internal olefins.; Knifton, John F.; J. Mol. Catal. (1987), 43(1), 65-77
- 7) Homogeneous cobalt, rhodium, and ruthenium catalysis adapted to the derivatization of allylic substrates with synthesis gas.; Lin, J. J.; Larkin, J. M.; Knifton, J. F.; New J. Chem. (1988), 12(6-7), 669-73
- 8) Process for 1-step preparation of bicyclic diamines as urethane catalysts.; Alexander, David C.; US 4705897
- 9) Syngas reactions. XII. The selective preparation of acetaldehyde, alcohols, esters and acetic acid from synthesis gas.; Knifton, John F.; Alexander, David C.; Isr. J. Chem. (1987), 1986, 27(3), 255-61
- 10) The selective preparation of ethylene glycol, alkanols, and acetic acid directly from synthesis gas.; Knifton, John F.; Int. Congr. Catal., [Proc.], 8th (1984), Volume 5, V39-V50
- 11) The selective generation of acetic acid directly from synthesis gas.; Knifton, John F.; ACS Symp. Ser. (1987), 328(Ind. Chem. C1 Processes), 98-107
- 12) C1-4-alkanol production from synthesis gas using a trimetallic catalyst; Knifton, John F., Grice, Neil J.; US 4622343
- 13) C1-C4 alcohol production from synthesis gas.; Knifton, John F.; US 4605677
- 14) Ethylene glycol and lower monohydric alcohols from syngas using a novel catalyst system; Grigsby, Robert A., Jr.; Knifton, John F.; US 4558072
- 15) Synthesizing formamides from synthesis gas plus ammonia.; Knifton, John F.; US 4556734
- 16) The selective generation of acetic acid directly from synthesis gas.; Knifton, John F.; Prepr. - Am. Chem. Soc., Div. Pet. Chem. (1986), 31(1), 26-31
- 17) Selective preparation of secondary and tertiary amines.; Knifton, John F., Alexander, David C.; US 4534411, EP 145191
- 18) Syngas reactions. IX. Acetic acid from synthesis gas.; Knifton, John F.; J. Catal. (1985), 96(2), 439-53
- 19) Acetaldehyde production from synthesis gas.; Lin, Jiang Jen; Knifton, John F.; US 4525481
- 20) Selectively preparing acetic anhydride by carbonylation of methyl acetate in the presence of an iodide-free catalyst system; Lin, Jiang Jen; Duranleau, Roger G.; US 4519956

- 21) Ethylene glycol using a ruthenium-copper catalyst.; Knifton, John F.; US 4518715
- 22) Methyl acetate from methanol and carbon monoxide using a catalyst system.;
Lin, Jiang Jen; US 4484002
- 23) Alcohols and aldehydes by reacting olefins with carbon monoxide and hydrogen.; Knifton, John Frederick; Lin, Jiang Jen; Grigsby, Robert Allison, Jr.; Brader, Walter Howe, Jr.; EP 107430
- 24) Mechanisms for the generation of alcohols, carboxylic acids and their esters directly from syngas.; Knifton, John F.; Prepr. - Am. Chem. Soc., Div. Pet. Chem. (1984), 29(2), 586-93
- 25) Alcohol/ester fuels from synthesis gas.; Knifton, John F.; Grigsby, Robert A., Jr.; Herbstman, Sheldon; Catal. Convers. Synth. Gas Alcohols Chem., [Proc. Symp.] (1984), 1983, 81-96.
- 26) Alcohols and esters from synthesis gas.; Lin, Jiang Jen; US 4436838
- 27) Alcohols from synthesis gas.; Lin, Jiang Jen; US 4436837, US 4434248, US 4362821
- 28) Acetaldehyde from methanol and synthesis gas using a novel catalyst composition.; Lin, Jiang Jen; Knifton, John F.; US 4433177, US 4433176, US 4433178
- 29) Homologation of methanol to ethanol.; Lin, Jiang Jen; Knifton, John F.; US 4424384, EP 56679
- 30) Syngas reactions. 6. Aliphatic alcohols and esters from synthesis gas.; Knifton, John F.; Grigsby, Robert A., Jr.; Lin, J. J.; Knifton, John F.; Grigsby, Robert A., Jr.; Lin, J. J.
- 31) Alkyl esters of carboxylic acids from an acid and syngas using a novel catalyst composition.; Lin, Jiang Jen; Knifton, John Frederick; GB 2113206
- 32) Ethylene glycol and lower monovalent alcohols.; Simons, Leslie Harold; DE 3243385, DE 3243386
- 33) Syngas reactions. V. Ethylene from synthesis gas.; Knifton, John F.; J. Catal. (1983), 79(1), 147-55
- 34) Acetic and propionic acids and their esters.; Knifton, John F.; US 4362822, US 4366259
- 35) Alcohols from synthesis gas.; Knifton, John F.; US 4339545, US 4332914
- 36) Alcohols from synthesis gas.; Knifton, John F.; Lin, Jiang Jen; US 4332915
- 37) Glycols and ethers.; Knifton, John F.; US 4315994
- 38) Ethylene from synthesis gas.; Knifton, John F.; US 4270015
- 39) "Use of an ionic composition, particularly an ionic liquid solvent such as an imidazolium salt, or a fused phosphonium salt, as a reagent for substitution reactions, a composition constituting a fluorination reagent, and a method of using it, e.g., for fluorination of aromatic and benzylic chlorides"; Garayt, Maxime; Le Boulaire, Virginie; Gree, Danielle; Gree, Rene; Schanen, Vincent; Spindler, Jean-Francis; WO 2002092608 ; FR 2824831
- 40) "A powerful, practical and chemoselective synthesis of 2-alkylalkanols catalyzed by $\text{Bi}(\text{TFA})_3$ or $\text{Bi}(\text{Otf})_3$ in the presence of molten TBAB"; M. M. Khodaei, A. R. Kosropour and K. Ghozati; Tetrahedron Letters, 2004, 45, 3525-3529.
- 41) "Zn/CeCl₃.&H₂O-TBPB: A New and "Green" Promoter System for Rapid and Regioselective Thiolyzation of 1,2-Epoxides with Aryl Disulfides"; A. R. Khosropour, M. M. Khodaei and K. Ghozati; Chemical Letters, 2004, 33, 1378-1379.
- 42) "The first platinum-catalyzed hydroamination of ethylene"; J J Brunet, M Cadena, N C Chu, O Diallo, K Jacob and E Mothes; Organometallics 2004, 23, 1264-1268

Important Notice: The information and statements in this data sheet are believed to be reliable but are not to be construed as a warranty or representation for which we assume legal responsibility or as an assumption of a duty on our part. Users should undertake sufficient verification and testing to determine the suitability for their own particular purpose of and information or products or vendors referred to within. **NO WARRANTY OF FITNESS FOR A PARTICULAR PURPOSE IS MADE.** Nothing in this data sheet is to be taken as permission, inducement or recommendation to practice any patented invention without a license.

Trademark Notice: The ® indicates a Registered Trademark in the United States and the TM or * indicates a Trademark in the United States. The mark may also be registered, the subject of an application for registration or a trademark in other countries.

Last updated: September 2, 2005