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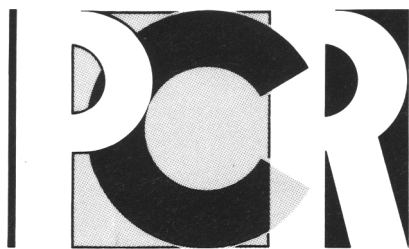
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Europe Langen, Germany Tel: 49 (0) 6103 708 301 Fax: 49 (0) 6103 708 310
Japan Tokyo, Japan Tel: (0473) 80-8500 Fax: (0473) 80-8505
Latin America Mexico City, Mexico Tel: 52-5-651-7077 Fax: 52-5-593-6223
Australia Melbourne, Australia Tel: (03) 212-8585 Fax: (03) 212-8502

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