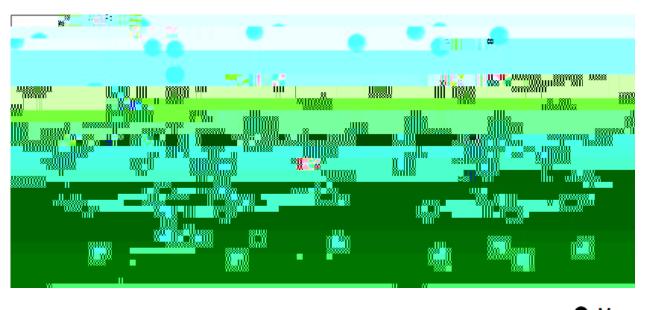
Transition-metal cyanides

This area of research has recently become very topical; many groups throughout the world are exploiting the fact that cyanide units can be linked together to form novel framework materials. These materials are of interest in a number of fields including supermolecular chemistry, hydrogen storage and crystal engineering. They show curious physical properties such as negative thermal expansion (i.e., they shrink when heated), negative compressability under pressure and luminescence ó all of which are exciting to study!

We employ chemical design to assemble one- two- and three-dimensional cyanide structures.

One-dimensional chain structures:

Although these have simple formulae, they can be structurally complex. For example, the group 11 cyanides, high-temperature CuCN,¹ AgCN² and AuCN³ are all constructed from linear metal-cyanide chains, [M E P _n packed together. They are highly disordered crystalline solids and their structure solution has involved the use of both X-ray and neutron Bragg and total scattering experiments.

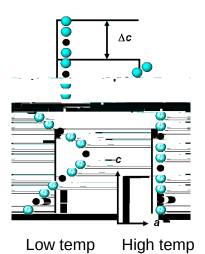


AgCN and HT- CuCN



● M ● C/N

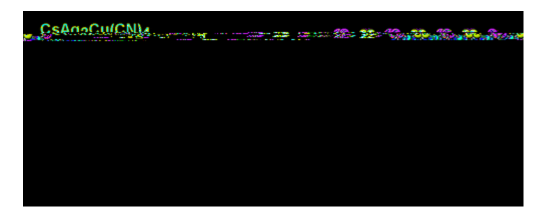
On heating these materials, they contract along the chain axis (c direction) i.e. show 1-D negative thermal expansion.⁴ A simple model to account for this behaviour is shown on the right. As the temperature is increased, there is an increase in amplitude of the transverse motions perpendicular to the chain axis leading to a kinking of the chains and a decrease in length along c.



Mixed-metal compounds such as $Cu_{1/2}Au_{1/2}(CN)$ and $Ag_{1/2}Au_{1/2}(CN)$ can be prepared⁵ enabling

Three-dimensional framework structures:

Dg{qpf vjgug ÷ukorngø ocvgtkenu vjg itqwr jeu dggp cevkxg kp rtgretkpi oqtg eqornkeevgf e{epkfg frameworks which demonstrate interesting stuetural features such as topological isomerism, polymorphism, and interpenetration, in which copies of the framework nest inside each other.¹⁰⁻¹² These materials can be prepared by solvothermal synthesis. The products are highly crystalline and their structures can be solved using single-crystal X-ray diffraction.



Above is shown the astonishing conversion between the two polymorphs of $CsAg_2Cu(CN)_4$. This conversion, which involves a major change in connectivity and topology, occurs at 180 °C as a single-crystal to single-crystal transformation.

Key References

30 (c)õEwEP< C rqn{oqtrjke ocvgtkcn0 Uvtwevwtg qh qpg hqto fgvgtokpgf htqo vqvcn pgwvtqp fkhhtcevkqp.ö U0L0 Jkddng. U0O0 Ejg{pg. C0E0 Jcppqp cpf U0I0 Gxgtuhkgnf. *Inorg. Chem.*, (2002), **41**, 66;20 (d) õ 340 õJgnkegu. Ejktcnkv{ cpf Kpvgtrgpgvtcvkqp< vjg Xgtucvknkty and Remarkable Interconversion of Silver-Eqrrgt E{cpkfg Htcogyqtmuö. C0O0 Ejkrrkpfcng cpf U0L0 Jkddng. *J.A.C.S.*, <u>131</u>, 127366 12744 (2009).